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05mar03 11:18:55 User208669 Session D2223.1

\$0.35 0.101 DialUnits File1

\$0.35 Estimated cost File1

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\$0.35 Estimated total session cost 0.101 DialUnits

File 155:MEDLINE(R) 1966-2003/Mar W1

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Set Items Description

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SI 8094 PARVO? OR AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT?

46011 TERMINUS OR TERMINI

3 141 SI AND S2

4 27843 TRANSPOS?

TOWNER CO.

1 S3 AND S4 342515 LEFT OR RIGHT

, 22 S3 AND S6

8 9668 TELOMER? 9 8 S1 AND S8 NOT S3

S10 74 BOVINE(W)PARVO?

? t s5/7/1

5/7/1

DIALOG(R)File 155:MEDLINE(R)

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Rescue of adeno-associated virus from recombinant plasmids: gene correction within the terminal repeats of AAV.

Samulski R J; Srivastava A; Berns K I; Muzyczka N

Cell (UNITED STATES) May 1983, 33 (1) p135-43, ISSN 0092-8674 fournal Code: 0413066

Contract/Grant No.: 5 R01 A116326; AI; NIAID; 5 T32 A107110; AI; NIAID

Document type: Journal Article

Languages: ENGLISH Main Citation Owner: NLM

Record type: Completed

We have isolated three types of pBR322-AAV recombinant plasmids that contain deletions within the 145 bp AAV terminal repeats. When the plasmids were transfected into human cells, mutants that contained deletions within the left (type I) or right (type II) terminal repeat were viable. Of four mutants examined that contained deletions in both termini (type III), only one was viable. All of the viable mutants produced AAV virions that contained wild-type AAV DNA. Furthermore, the viable type III deletion

could be converted to a nonviable mutant by deleting all copies of an 11 bp sequence from its termini. We conclude that there is an efficient mechanism for correcting deletions within the AAV termini. A model that could account for these observations is also discussed.

Record Date Created: 19840928

? t s5/5/1

1/5/1

DIALOG(R)File 155:MEDLINE(R)

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Rescue of adeno-associated virus from recombinant plasmids: gene correction within the terminal repeats of AAV.

Samulski R J; Srivastava A; Berns K I; Muzyczka N

Cell (UNITED STATES) May 1983, 33 (1) p135-43, ISSN 0092-8674

Iournal Code: 0413066

Contract/Grant No.: 5 R01 AI16326; AI; NIAID; 5 T32 AI07110; AI; NIAID

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

Subfile: INDEX MEDICUS

We have isolated three types of pBR322-AAV recombinant plasmids that contain deletions within the 145 bp AAV terminal repeats. When the plasmids were transfected into human cells, mutants that contained deletions within the 1eft (type I) or right (type II) terminal repeat were viable. Of four mutants examined that contained deletions in both termini (type III), only one was viable. All of the viable mutants produced AAV virions that contained wild-type AAV DNA. Furthermore, the viable type III deletion could be converted to a nonviable mutant by deleting all copies of an 11 bp sequence from its termini. We conclude that there is an efficient mechanism for correcting deletions within the AAV termini. A model that could account for these observations is also discussed.

Tags: Human; Support, Non-U.S. Gov't; Support, U.S. Gov't, P.H.S.

Descriptors: *Dependovirus--genetics--GE; *Virus Replication; Chromosome

Deletion, DNA Replication, DNA Transposable Elements, DNA, Recombinant, DNA, Viral--genetics--GE, Defective Viruses--genetics--GE, Mutation;

Plasmids; Repetitive Sequences, Nucleic Acid

Molecular Sequence Databank No: GENBANK/J01901; GENBANK/M12405,

CAS Registry No.: 0 (DNA Transposable Elements); 0 (DNA, Recombinant)

GENBANK/M12468; GENBANK/M12469

Record Date Created: 19840928

0 (DNA, Viral); 0 (Plasmids)

7 t s7/7/7

11111

DIALOG(R)File 155:MEDLINE(R)

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Interaction of virally coded protein and a cell cycle-regulated cellular protein with the bovine parvovirus left terminus ori.

Metcalf J B; Bates R C; Lederman M

Biology Department, Virginia Polytechnic Institute and State University,

3lacksburg 24061-0406.

Journal of virology (UNITED STATES) Nov 1990, 64 (11) p5485-90,

SSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

conformation) of BPV. Three specific DNA-protein complexes formed. One complex was shown to involve a BPV structural protein(s) by inhibiting its formation when antiserum specific for these BPV proteins was used. By nonstructural proteins, a second complex was shown to involve a BPV specific competition with serum containing antibodies against the BPV DNA competition assays suggest that the viral proteins do not bind to the contact-inhibited cells. Since autonomous parvovirus replication requires nonstructural protein. A third complex contained protein of cellular origin right hairpin, which differs in sequence and secondary structure from the an S-phase factor for progeny formation, the terminal binding protein trans-acting viral and cellular proteins. A gel retardation assay was used (BPV)-infected bovine fetal lung cells that interact with the hairpinned and was also formed with extracts of uninfected bovine fetal lung cells. palindromes that function as origins of replication in conjunction with preventing formation of the stem of the hairpin. The cellular protein is Replication of parvoviruses requires cis signals located in terminal left terminus, or to a BPV terminus that lacks the first 52 nucleotides, to identify proteins in crude nuclear extracts of bovine parvovirus regulated in a cell cycle-dependent fashion, with its binding activity increased in uninfected, actively dividing cells compared with left end (3' OH terminus of the viral minus strand in the flop demonstrated here is a candidate for this factor.

Record Date Created: 19901115

?ts7772312131617

DIALOG(R)File 155:MEDLINE(R)

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.0378075 99355161 PMID: 10428207

cis requirements for the efficient production of recombinant DNA vectors based on autonomous parvoviruses.

Kestler J; Neeb B; Struyf S; Van Damme J; Cotmore SF; D'Abramo A; Tattersall P; Rommelaere J; Dinsart C; Cornelis J J

Applied Tumor Virology Abt. F0100 and INSERM U375 Deutsches Krebsforschungszentrum, Heidelberg, Germany.

Human gene therapy (UNITED STATES) Jul 1 1999, 10 (10) p1619-32, SSN 1043-0342 Journal Code: 9008950

Contract/Grant No.: CA29303; CA; NCI Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM Record type: Completed

vectors from infectious molecular clones of parvoviruses MVMp and H1 were vectors from improved recombinant molecular DNA clones amounted to 5 x genome, with a mere 6% increase in DNA length leading to an approximately were reduced more than 50 times in comparison with recombinant vectors in equivalent-length sequence consisting of reporter cDNA and stuffer DNA, genomes into recombinant particles appeared to depend on in cis-provided 800 bp was removed from the VP transcription unit. Furthermore, titers of viral vectors, in which most of the VP-coding region was replaced by an which stuffer DNA was not substituted for the residual VP sequence. In allow the assessment of the therapeutic effect of recombinant parvoviruses The replication of viral genomes and the production of recombinant viral cotransfection of recombinant clones and helper plasmids providing the greatly improved by the introduction of a consensus NS-1 nick site at the unction between the left-hand viral terminus and the plasmid DNA. structural genes as well as insertions of foreign DNA in replacement of recombinant H1 virus genomes. In contrast, the incorporation of these structural gene sequences. Indeed, the production of H1 viral vectors by structural proteins (VPs) in trans, drastically decreased when more than above-mentioned requirements for efficient packaging, titers of virus Progressive deletions of up to 1600 bp in the region encoding the those sequences did not appreciably affect the replication ability of the addition, viral vector production was restricted by the overall size of the 0 times lower encapsidation yield. Under conditions fulfilling the 10(7) infectious units per milliliter of crude extract. These titers should expressing small transgenes in laboratory animals.

Record Date Created: 19990922

DIALOG(R)File 155:MEDLINE(R)

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The minute virus of mice (MVM) nonstructural protein NS1 induces nicking of MVM DNA at a unique site of the right-end telomere in both hairpin and duplex conformations in vitro.

Willwand K; Baldauf A Q; Deleu L; Mumtsidu E; Costello E; Beard P; Rommelaere J

and Formation INSERM U375, Heidelberg, Germany. k.willwand@dkfz-heidelberg. Deutsches Krebsforschungszentrum, Department of Applied Tumor Virology,

Oct 1997, 78 (Pt 10) p2647-55, Journal of general virology (ENGLAND) ISSN 0022-1317 Journal Code: 0077340

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

The right-end telomere of replicative form (RF) DNA of the autonomous extract supplemented with the MVM nonstructural protein NS1. This extended form, base-paired to a copy strand (duplex conformation). We processing is shown here to result from the NSI-dependent nicking of the complementary strand at a unique position 21 nt inboard of the folded-back end of the template strand, possibly due to NS1 which is covalently bound symmetry and an interior bulge of three unpaired nucleotides on one strand parvovirus minute virus of mice (MVM) consists of a sequence that is self-complementary except for a three nucleotide loop around the axis of conformation, while processing of the duplex structure leads to the release to this end. A fraction of the right-end duplex product undergoes melting synthesis at the right terminus stops a few nucleotides before reaching the exist in the form of a folded-back strand (hairpin conformation) or in an of free right-end telomeres. In the majority of molecules, displacement recently reported that the right-end telomere is processed in an A9 cell distinguishing these structures are not prerequisites for nicking under the (designated the right-end 'bubble'). This right-end inverted repeat can genomic 5' end. DNA species terminating in duplex or hairpin by strand-displacement synthesis, generating the right-end duplex bulge, are all cleaved in the presence of NS1, indicating that features in vitro conditions tested. Cleavage of the hairpin structure is followed configurations, or in a mutated structure that has lost the right-end and re-folding into hairpin structures (formation of a 'rabbit-ear'

Record Date Created: 19971117

71///

DIALOG(R)File 155:MEDLINE(R)

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Interactions between the termini of adeno-associated virus DNA.

Bohenzky R A; Berns K I

Department of Immunology and Medical Microbiology, College of Medicine, Jaiversity of Florida, Gainesville 32610.

Journal of molecular biology (ENGLAND) Mar 5 1989, 206 (1) p91-100,

Contract/Grant No.: AI-01770; AI; NIAID; AI-22251; AI; NIAID

SSN 0022-2836 Journal Code: 2985088R Contract/Grant No. ALO1770, AT MIAID

Languages: ENGLISH

Document type: Journal Article

Main Citation Owner: NLM

Record type: Completed

The adeno-associated virus (AAV) genome is a linear, single polynucleotide chain with inverted terminal repeats of 145 bases. In order to test whether the terminal repeats at opposite ends of the genome have to

chimeric genomes in which an 11 base symmetrical sequence has been deleted When plasmid clones were used, the structure of virion DNA depended on the passages contained DNA with wild-type sequences in both terminal repeats. DNA to transfect adenovirus-infected HeLa cells. When chimeric duplex be able to completely base-pair during DNA replication, we have created opposite ends of the genome may interact during DNA replication, it is not different 12 base symmetrical sequence. We have used these chimeric repeat sequences were again wild-type. However, if the original construct the wild-type sequence displays an advantage over the mutant allele. (3) In constructs either as a duplex insert in pBR322 or as purified duplex virion necessary that they be perfectly complementary. (2) In direct competition, right end of the genome (terminus of genetic map), all progeny terminal virion DNA was used, all of the progeny virions obtained after two cell progeny molecules were parental in type (i.e. mutant left and wild-type right terminal repeat). We conclude (1) although the terminal repeats at a plasmid clone, the terminal repeat on the left end of the genome is at an from the terminal repeat at one end of the genome and replaced by a contained the mutant sequence in the left terminal repeat, the majority of advantage in a competitive situation. We note that the left terminal repeat original orientation. If the mutant terminal repeat was originally at the is adjacent to a transcriptional promoter.

Record Date Created: 19890525

7/13

DIALOG(R)File 155:MEDLINE(R)

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Analysis of the termini of the DNA of bovine parvovirus: demonstration of sequence inversion at the left terminus and its implication for the replication model.

Chen K C; Shull B C; Lederman M; Stout E R; Bates R C

Department of Biology, Virginia Polytechnic Institute and State University, Blacksburg 24061.

Journal of virology (UNITED STATES) Oct 1988, 62 (10) p3807-13,

ISSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

The distribution of terminal-sequence orientations in the viral DNA of bovine parvovirus (BPV), an autonomous parvovirus, was studied by end labeling and restriction enzyme digestion and also by cloning. The left (3') end of the minus strand of BPV was found in two alternative sequence orientations (designated as flip and flop, which are reverse complements of each other), with a 10-fold excess of flip. This is in contrast to the autonomous rodent parvoviruses which encapsidate minus-strand DNA with only the flip orientation at this end. The right (5') end of the minus strand of

the plus strand, which makes up about 10% of the encapsidated BPV DNA. Each parvoviruses, cannot account for the observed distribution of BPV DNA. An Replicative-form DNA showed the same distribution of terminal-sequence rodent parvoviruses. Sequence inversions were also detected at both ends of orientations as the reannealed plus and minus virion DNAs, suggesting that BPV contained both sequence orientations with equal frequencies, as in the this ratio was restored in the progeny DNA resulting from transfection with terminus of BPV DNA had a characteristic ratio of flip to flop forms, and he distribution of flip and flop forms observed in virion DNA is not due replicative forms. The current replication model for autonomous parvoviruses, which was based on the available data for the rodent genomic clones of different defined terminal conformations. to selective encapsidation, but rather to the specific distribution of alternative model is suggested.

Record Date Created: 19881019

DIALOG(R)File 155:MEDLINE(R)

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minute virus of mice DNA suggests a modified rolling hairpin model for Sequence analysis of the termini of virion and replicative forms of autonomous parvovirus DNA replication.

Astell CR; Chow MB; Ward DC

Journal of virology (UNITED STATES) Apr 1985, 54 (1) p171-7, ISSN

0022-538X Journal Code: 0113724

Contract/Grant No.: CA-16038; CA; NCI; GM-20124; GM; NIGMS

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

pairs beyond that expected from the known sequence of the virion DNA. These (5') terminus was sequence heterogeneous and extended an additional 18 base data unambiguously establish the sequence complexity at the termini of both comparison of the combined sequence information leads us to propose a of mice, were determined. The left (3') terminus had a unique sequence on The nucleotide sequences of the terminal regions of monomer replicative form DNA, a pivotal intermediate species in the replication of minute virus the single-stranded viral genome and the pool of replicative DNA. A both strands and in both 3'-hairpin configurations. In contrast, the right modified rolling hairpin model for the replication of autonomous parvoviruses which is compatible with all available data.

Record Date Created: 19850418

DIALOG(R)File 155:MEDLINE(R)

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Rescue of adeno-associated virus from recombinant plasmids: gene correction within the terminal repeats of AAV.

Samulski R J, Srivastava A; Berns K I, Muzyczka N

Cell (UNITED STATES) May 1983, 33 (1) p135-43, ISSN 0092-8674

Iournal Code: 0413066

Contract/Grant No.: 5 R01 AI16326; AI; NIAID; 5 T32 AI07110; AI; NIAID

Document type: Journal Article Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

contain deletions within the 145 bp AAV terminal repeats. When the plasmids We have isolated three types of pBR322-AAV recombinant plasmids that for correcting deletions within the AAV termini. A model that could account contained wild-type AAV DNA. Furthermore, the viable type III deletion sequence from its termini. We conclude that there is an efficient mechanism could be converted to a nonviable mutant by deleting all copies of an 11 bp were transfected into human cells, mutants that contained deletions within one was viable. All of the viable mutants produced AAV virions that mutants examined that contained deletions in both termini (type III), only the left (type I) or right (type II) terminal repeat were viable. Of four for these observations is also discussed.

Record Date Created: 19840928

?ts9/1/67

DIALOG(R)File 155:MEDLINE(R)

(c) format only 2003 The Dialog Corp. All rts. reserv.

Specific initiation of replication at the right-end telomere of the closed species of minute virus of mice replicative-form DNA.

Baldauf A Q; Willwand K; Mumtsidu E; Nuesch J P; Rommelaere J

Department of Applied Tumor Virology, Deutches Krebsforschungszentrum, Heidelberg, Germany.

Journal of virology (UNITED STATES) Feb 1997, 71 (2) p971-80, ISSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

(MVM). MVM virion DNA, a single-stranded molecule bracketed by short, natural DNA templates of the autonomous parvovirus minute virus of mice replicative-form (RF) DNA when incubated in mouse A9 fibroblast extract. covalently closed RF (cRF) molecule as the major conversion product. cRF erminal, self-complementary sequences, is converted into double-stranded The 3' end of the newly synthesized complementary strand is ligated to the We have developed an in vitro system that supports the replication of right-end hairpin of the virion strand, resulting in the formation of a

by immunoprecipitation with NS1-specific antibodies. The 5'eRF product is ourified MVM nonstructural protein NS1 expressed from recombinant covalently attached to the right-end telomere of the DNA product, as shown of current models of parvovirus DNA replication and provide new insights baculoviruses or vaccinia viruses, cRF DNA is processed into a right-end structure for recognition of the DNA substrate by NS1. Further processing he target for additional rounds of NS1-induced nicking and displacement telomere. Formation of dRF DNA is highly stimulated by NS1. The followed by unfolding of the hairpin and copying of the terminal sequence. presence of NS1. In the course of the right-end nicking reaction, NS1 gets experimental results presented in this report support various assumptions DNA in a left-to-left-end configuration, presumably as a result of copying of the whole molecule by displacement synthesis initiated at the right-end of the 5'eRF template in vitro leads to the formation of dimeric RF (dRF) ONA is not further replicated in A9 cell extract alone. On addition of synthesis at the right end, arguing against the requirement of the hairpin (5' end of the virion strand) extended form (5'eRF). This is indicative of NS1-dependent nicking of the right-end hairpin at a distinct position, in contrast, no resolution of the left-end hairpin can be detected in the nto the replication functions of the NS1 protein.

Record Date Created: 19970218

DIALOG(R)File 155:MEDLINE(R)

(c) format only 2003 The Dialog Corp. All rts. reserv.

An asymmetric nucleotide in the parvoviral 3' hairpin directs segregation of a single active origin of DNA replication.

Cotmore S F; Tattersall P

Department of Laboratory Medicine, Yale University School of Medicine,

EMBO journal (ENGLAND) Sep 1 1994, 13 (17) p4145-52, ISSN 0261-4189 New Haven, CT 06510.

Contract/Grant No.: AI26109; AI; NIAID; CA29303; CA; NCJ

ournal Code: 8208664

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

of mice (MVM), a murine parvovirus, can assume a complex hairpin structure. copied to form an imperfect palindrome which bridges adjacent genomes in a This contains a stem in which there is a mismatched 'bubble' sequence where dimer duplex intermediate, leaving the two 'bubble' sequences embedded in The 3' telomere of the linear single-stranded DNA genome of minute virus unctions are resolved asymmetrically in vitro in a DNA synthetic reaction which requires the viral initiator protein NS1. We show that the sequence a GA doublet opposes a GAA triplet. During replication, this hairpin is otential replication origins on either side of the axis of symmetry. Such

approximately 50 bp long, extending from an Activated Transcription Factor ounding site at one end to a position some 7 bp beyond the major initiation site, to which NS1 ultimately becomes covalently attached. The actual critical spacer element. Segregation of this asymmetry, therefore, allows sequence of the GA doublet is unimportant, but insertion of any third surrounding the doublet is a potent origin, but the analogous region nucleotide here mactivates the origin, indicating that it represents a he virus to confine replication initiation to one particular telomeric containing the triplet is completely inactive. The active origin is configuration.

Record Date Created: 19941005

? s bovine(w)parvo?

122385 BOVINE

6618 PARVO?

S10 74 BOVINE(W)PARVO?

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Temp SearchSave "TD794" stored

05mar03 11:31:17 User208669 Session D2223.2

\$6.62 2.069 DialUnits File155 \$0.21 1 Type(s) in Format 5

\$0.00 105 Type(s) in Format 6 \$2.10 10 Type(s) in Format 7

\$2.31 116 Types

\$8.93 Estimated cost File155

\$3.02 TELNET

\$11.95 Estimated cost this search

\$12.30 Estimated total session cost 2.170 DialUnits

File 50:CAB Abstracts 1972-2003/Jan

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*File 50: Truncating CC codes is recommended for full retrieval See Help News50 for details.

Set Items Description

? s bovine (w)parvo?

75866 BOVINE 4122 PARVO?

176 BOVINE (W)PARVO?

? s vector? and s1

54093 VECTOR? 176 SI

1 VECTOR? AND S1

? t s2/6

00962529 CAB Accession Number: 802261627

Document Type: Journal article ISSN: 0012-4966 1122 AAV Department of Biological Sciences, Purdue University, West Lafayette, Structure, sequence, and function correlations among parvoviruses. *File 50: Truncating CC codes is recommended for full retrieval. \$13.92 Estimated total session cost 2.481 DialUnits 05mar03 11:31:52 User208669 Session D2223.3 Tropical veterinary science. Research report 1979. 33013408 CAB Accession Number: 952206035 Virology (New York) vol. 194 (2): p.491-508 File 155:MEDLINE(R) 1966-2003/Mar W1 (c) format only 2003 The Dialog Corp. c) 2003 CAB International. All rts. reserv. File 50:CAB Abstracts 1972-2003/Jan \$0.00 1 Type(s) in Format 6 \$1.40 0.311 DialUnits File50 SYSTEM:OS - DIALOG OneSearch S1 250 BOVINE(W)PARVO? DIALOG(R)File 50:CAB Abstracts Chapman, M. S.; Rossmann, M. G. \$1.62 Estimated cost this search (c) 2003 CAB International S2 194 RD (unique items) 2/7/85 (Item 11 from file: 50) \$1.40 Estimated cost File50 ..completed examining records ..examined 50 records (100) ..examined 50 records (150) ..examined 50 records (200) ...examined 50 records (250) See Help News50 for details. ...examined 50 records (50) Set Items Description Publication Year: 1979 Publication Year: 1993 198251 BOVINE 10740 PARVO? \$0.00 1 Types Indiana 47907, USA. ISSN: 0042-6822 --? s bovine(w)parvo? anguage: English \$0.22 TELNET ? t s2/7/85 91

MDV DNA with the AUG codon of the 1st ORF underlined, and the region of including human parvovirus (HPV), adeno-associated virus 2 (AAV2), bovine virus (densovirus) of mosquitoes (MDV), a member of the Parvoviridae which parvo)virus of mice (MVM). Studies are underway to determine the complete parvovirus (BVP), Aleutian disease (parvo)virus of mink (ADV) and minute the sequence of the DNA "plus" chain and 3 extended open reading frames Translated from Doklady Akademii Nauk SSSR, 307: 996-1000 (1989) -preparation Viroden has been developed from it. The present paper shows greatest homology of the NS1 proteins of MDV and other parvoviruses (ORFs), with the probable promoters and polyadenylation signals for these V.A. Engel'gardt Institute of Molecular Biology, Academy of Sciences of ORFs underlined, the 3'-terminal sequence of the virion "minus" chain of Galev, E. E.; Afanas'ev, B. N.; Buchatskii, L. P.; Kozlov, Yu. V.; Baev, The nucleotide sequence is given of the virion of the densonucleosis includes the smallest known viruses pathogenic to insects. This virus infects the larvae of Culex, Aedes and Culiseta spp. and the insecticide Features of organization of the genome of the densoviruses. Doklady, Biological Sciences vol. 307 (1-6): p.537-541 ? s parvo? or aav or adenoassociat? or adeno(w)associat? 32467824 CAB Accession Number: 910504273 S3 4946 ORIGIN(2N)REPLICATION (c) 2003 CAB International. All rts. reserv. 1647 ADENO(W)ASSOCIAT? Publication Year: 1989, publ. 1990 the USSR, Moscow, RSFSR, USSR. 250 BOVINE(W)PARVO? **DIALOG(R)File 50:CAB Abstracts** primary structure of MDV. 15 ref. Document Type: Journal article 43 ADENOASSOCIAT? (Item 17 from file: 50) 194 RD (unique items) 108039 REPLICATION 1384056 ASSOCIAT? Items Description ? s origin(2n)replication 188736 ORIGIN 10740 PARVO? 2729 ADENO Language: English

S4 12290 PARVO? OR AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT? 12290 S4 4946 S3 ? s s3 and s4

59 S3 AND S4 SS

.. completed examining records ...examined 50 records (50)

S6 55 RD (unique items)

? t s6/7/8 9 20 32 35 38 48 50 51 52 6/7/8 (Item 8 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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Adeno-associated virus (AAV) site-specific recombination does not require a Rep-dependent origin of replication within the AAV terminal repeat. Young S M; Samulski R J

Curriculum in Genetics and Molecular Biology, Gene Therapy Center, Jniversity of North Carolina, Chapel Hill, NC 27599, USA.

Proceedings of the National Academy of Sciences of the United States of

America (United States) Nov 20 2001, 98 (24) p13525-30, ISSN 3027-8424 Journal Code: 7505876

Contract/Grant No.: DK51880; DK; NIDDK; HL 48347; HL; NHLBI

Document type: Journal Article

Main Citation Owner: NLM Languages: ENGLISH

Record type: Completed

required for AAV DNA replication are located on chromosome (ch) 19. Both extensive rearrangement and amplification of ch-19 sequences independent of of targeted integration in human cells. An AAV Rep binding element (RBE) Adeno-associated virus (AAV) is the only known eukaryotic virus capable terminal repeat (ITR) cis-acting sequences in targeted integration an AAV cotransfection in the presence or absence of Rep. Our results demonstrated genomes were unable to rescue and replicate. In addition, Rep78 induced similar frequency. Molecular characterization of the mutant ITR integrants prerequisite for site-specific recombination and suggests AAV targeting is tested. Complementation analysis confirmed that the mutant targeted viral irs mutant incapable of supporting viral replication was tested. Wild-type that, in the presence of Rep78, both ITR substrates targeted to ch-19 with ch-19 RBE and trs elements have been shown to be essential for viral wild-type or mutant targeting substrate. These studies demonstrate that confirmed the presence of the trs mutation in the majority of samples targeting to this locus. To characterize the role of the AAV inverted Rep-dependent nicking of the viral cis-acting trs sequence is not a and terminal resolution site (trs) identical to the viral terminal repeats and mutant substrates were assayed for targeted integration after

understanding of AAV site-specific recombination and the development of mediated by Rep78/68-dependent replication from the ch-19 origin of replication (ori). These studies have significant impact toward the targeting vectors.

Record Date Created: 20011121

6/7/9 (Item 9 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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transcription factor must form a precise ternary complex with origin DNA Minute virus of mice initiator protein NS1 and a host KDWK family for nicking to occur.

Christensen J; Cotmore S F; Tattersall P

Institute of Medical Microbiology and Immunology, University of

Copenhagen, Panum Institute, Copenhagen 2200 N, Denmark.

Journal of virology (United States) Aug 2001, 75 (15) p7009-17,

SSN 0022-538X Journal Code: 0113724

Contract/Grant No.: AI26109; AI; NIAID

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

preferentially enhanced on oriL(TC) compared to oriL(GAA). Without ATP, NS1 can be flexibly spaced. When coexpressed from recombinant baculoviruses, subunits of PIF, p79 and p96, cooperatively bind two ACGT half-sites, which concatemers whose junctions are resolved to give unit-length genomes by a nucleotide inserted between the NS1 and PIF sites. Here we examined the the distance between the NS1 binding site and the NS1-proximal half-site is half-site is unimportant. When expressed separately, both PIF subunits form he PIF subunits preferentially form heterodimers which, in the presence of interactions on oriL(TC) which lead to activation of NS1 by PIF. The two ATP, show cooperative binding with NS1 on oriL, but this interaction is process involving DNA replication initiated at origins derived from each viral telomere. The left-end origin of minute virus of mice (MVM), oriL, initiation factor (PIF), a member of the emerging KDWK family of Parvoviral rolling hairpin replication generates palindromic genomic contains binding sites for the viral initiator nickase, NS1, and parvovirus resistant to DNase I. Varying the spacing of the PIF half-sites shows that ranscription factors. oriL is generated as an active form, oriL(TC), and as an inactive form, oriL(GAA), which contains a single additional homodimers that bind site specifically to oril, but only complexes nteraction, rendering the NS1 binding site, but not the nick site, critical for nickase activation, whereas the position of the distal is unable to bind stably to its cognate site, but PIF facilitates this containing p79 activate the NS1 nickase function.

Record Date Created: 20010703

677/20 (Item 20 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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Packaging cells based on inducible gene amplification for the production of adeno-associated virus vectors.

Inoue N; Russell D W

Markey Molecular Medicine Center and Department of Medicine, University

Journal of virology (UNITED STATES) Sep 1998, 72 (9) p7024-31, of Washington, Seattle, Washington 98195, USA.

SSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

Although vectors based on adeno-associated virus (AAV) offer several

unique advantages, their usage has been hampered by the difficulties

The packaging and producer cell lines developed express SV40 T antigen encountered in vector production. In this report, we describe a new AAV packaging system based on inducible amplification of integrated helper and vector constructs containing the simian virus 40 (SV40) replication origin.

with approximately 10(4) vector particles produced per cell. These stocks allows inducible amplification of chromosomal loci linked to the SV40 titers that were 10 times higher than those obtained by standard methods, adenovirus infection resulted in helper and vector gene amplification as under the control of the reverse tetracycline transactivator system, which origin. Culturing these cells in the presence of doxycycline followed by well as higher vector titers. Clonal producer cell lines generated vector

production of vector stocks for human gene therapy. Record Date Created: 19980916

transfection step combined with the reproducibility of stable producer

were free of detectable replication-competent virus. The lack of a

lines makes this packaging method ideally suited for the large-scale

DIALOG(R)File 155:MEDLINE(R) 6/7/32 (Item 32 from file: 155)

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Minimum origin requirements for linear duplex AAV DNA replication in

Ward P; Berns K I

Department of Microbiology, W. R. Hearst Microbiology Research Center,

Virology (UNITED STATES) Jun 1 1995, 209 (2) p692-5, ISSN 0042-6822 Cornell University Medical College, New York, New York 10021, USA. Iournal Code: 0110674

Contract/Grant No.: GM50023; GM; NIGMS

Document type: Journal Article

Main Citation Owner: NLM Languages: ENGLISH

Record type: Completed

virus (AAV) Rep 68/78 protein. When a linear duplex template was used, initiation of one round of DNA replication was achieved when nucleotides We have investigated the minimal requirements for a functional origin of DNA replication in an in vitro assay which requires the adeno-associated required. Deletion of an additional 11 nucleotides from the terminal repeat -56 of the AAV inverted terminal repeat (in the flop orientation) were present at the terminus of the template. A terminal resolution site was not plocked the reaction.

Record Date Created: 19950711

6/7/35 (Item 35 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

(c) format only 2003 The Dialog Corp. All rts. reserv.

An asymmetric nucleotide in the parvoviral 3' hairpin directs segregation of a single active origin of DNA replication.

Cotmore S F; Tattersall P

Department of Laboratory Medicine, Yale University School of Medicine,

New Haven, CT 06510.

EMBO journal (ENGLAND) Sep 1 1994, 13 (17) p4145-52, ISSN 0261-4189 lournal Code: 8208664

Contract/Grant No.: AI26109; AI; NIAID; CA29303; CA; NCI

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

of mice (MVM), a murine parvovirus, can assume a complex hairpin structure. copied to form an imperfect palindrome which bridges adjacent genomes in a This contains a stem in which there is a mismatched 'bubble' sequence where approximately 50 bp long, extending from an Activated Transcription Factor dimer duplex intermediate, leaving the two 'bubble' sequences embedded in The 3' telomere of the linear single-stranded DNA genome of minute virus junctions are resolved asymmetrically in vitro in a DNA synthetic reaction which requires the viral initiator protein NS1. We show that the sequence a GA doublet opposes a GAA triplet. During replication, this hairpin is binding site at one end to a position some 7 bp beyond the major initiation critical spacer element. Segregation of this asymmetry, therefore, allows site, to which NS1 ultimately becomes covalently attached. The actual potential replication origins on either side of the axis of symmetry. Such sequence of the GA doublet is unimportant, but insertion of any third surrounding the doublet is a potent origin, but the analogous region he virus to confine replication initiation to one particular telomeric nucleotide here inactivates the origin, indicating that it represents a containing the triplet is completely inactive. The active origin is

Record Date Created: 19941005 configuration.

OIALOG(R)File 155:MEDLINE(R) 6/7/38 (Item 38 from file: 155)

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Features of the adeno-associated virus origin involved in substrate ecognition by the viral Rep protein.

Snyder R O; Im D S; Ni T; Xiao X; Samulski R J; Muzyczka N

Department of Microbiology, State University of New York at Stony Brook Medical School 11794-8621 Journal of virology (UNITED STATES) Oct 1993, 67 (10) p6096-104,

SSN 0022-538X Journal Code: 0113724

Contract/Grant No.: 5 PO1 CA2814607; CA; NCI; AI25530; AI; NIAID; RO1

3M3572302; GM; NIGMS

Document type: Journal Article Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

We previously demonstrated that the adeno-associated virus (AAV) Rep68 and Rep78 proteins are able to nick the AAV origin of DNA replication at

the terminal resolution site (trs) in an ATP-dependent manner. Using four types of modified or mutant substrates, we now have investigated the substrate requirements of Rep68 in the trs endonuclease reaction. In the

internal palindromes of the AAV terminal repeat were active in the first kind of substrate, portions of the hairpinned AAV terminal repeat were deleted. Only deletions that retained virtually all of the small

biochemical evidence that the secondary structure of the terminal repeat was an important feature for substrate recognition. In the second type of endonuclease reaction. This result confirmed previous genetic and

genome. The mutant was nicked at a 50-fold-lower frequency relative to a substrate, the trs was moved eight bases further away from the end of the wild-type origin, and the nick occurred at the correct trs sequence despite

mismatch substrates contained a wild-type sequence on the strand normally mismatch mutants were capable of being nicked in the presence of ATP cut but an incorrect sequence on the complementary strand. All of the secondary structure. It also suggested that the minimum trs recognition sequence extended three bases from the cut site in the 3' direction. The third type of substrate harbored mismatched base pairs at the trs. The its new position. This finding indicated that the endonuclease reaction required a specific sequence at the trs in addition to the correct

suggesting that the sequence on the opposite strand may also be recognized during nicking. Analysis of the mismatch mutants also suggested that a nterpretation was confirmed by analysis of the fourth type of substrate single-stranded trs was a viable substrate for the enzyme. This However, there was substantial variation in the level of activity,

the enzyme. We concluded that substrate recognition by the AAV Rep protein nvolves at least two and possibly as many as four features of the AAV cleaved efficiently by the enzyme provided that the correct strand was wild-type did. This finding indicated that the sequence at the cut site was not involved in recognition of the terminal repeat for specific binding by ested, which contained a single-stranded trs. This substrate was also substrates with mutant trss bound the Rep protein as efficiently as the present in the substrate. In addition, the single-stranded substrate no onger required ATP as a cofactor for nicking. Finally, all of the terminal repeat (ABSTRACT TRUNCATED AT 400 WORDS)

Record Date Created: 19931012

DIALOG(R)File 155:MEDLINE(R) (Item 48 from file: 155)

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Replication of adeno-associated virus DNA. Complementation of naturally occurring rep- mutants by a wild-type genome or an ori- mutant and correction of terminal palindrome deletions.

Senapathy P; Tratschin JD; Carter BJ

Journal of molecular biology (ENGLAND) Oct 15 1984, 179 (1) p1-20,

SSN 0022-2836 Journal Code: 2985088R

Document type: Journal Article Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

on replication of AAV DNA. We obtained AAV mutants by molecular cloning in when the recombinant AAV-plasmid DNA is transfected into human 293 cells internal deletion of AAV coding sequences. Also, some of these mutant-AAV one terminal palindrome was phenotypically wild-type and allowed rescue and plasmids have additional deletions of one or both AAV terminal palindromes replication (rep-) but could be complemented by intact wild-type AAV. This introduced during constructions in vitro. We show here that AAV mutants indicates that an AAV replication function, Rep, is required for normal AAV replication of AAV genomes in which the deleted region was regenerated apparently by an intramolecular correction mechanism. One model for this bacterial plasmid, infectious AAV genomes can be rescued and replicated When the entire adeno-associated virus (AAV) genome is inserted into a were also replication defective but were not complementable by wild-type replication. Mutants in which both terminal palindromes were deleted (ori-) together with helper adenovirus particles. We have taken advantage of this AAV. The cis-dominance of the ori- mutation shows that the replication defective-interfering genomes. Each of these mutants contains a single containing internal deletions were defective for replicative form DNA experimental system to analyze the effects of several classes of mutations origin is comprised in part of the terminal palindrome. Deletion of only bacterial plasmids of naturally occurring AAV variant or

correction mechanism is proposed. An AAV ori- mutant also complemented mutants in that monomeric single-stranded single-stranded DNA accumulated replication of AAV rep- mutants as efficiently as did wild-type AAV. These very inefficiently even though monomeric single-stranded DNA from the studies also revealed an unexpected additional property of the deletion complementing wild-type AAV did accumulate.

Record Date Created: 19841228

677/50 (Item 50 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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DNA sequence of the 5' terminus containing the replication origin of parvovirus replicative form DNA.

Rhode S L; Klaassen B

Journal of virology (UNITED STATES) Mar 1982, 41 (3) p990-9, ISSN 0022-538X Journal Code: 0113724

Contract/Grant No.: CA-25866; CA; NCI; CA26801; CA; NCI

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

noncoding and contains a 55-base-pair tandem repeat. The addition mutant of H-1, DI-1, was also sequenced in this region and shown to have three copies palindrome in native replicative form DNA, one inverted with respect to the replication origin for parvovirus replicative form DNA replication. Some of The nucleotide sequence of the 5' terminus of the parvovirus H-1 was contains only one copy of this repeat sequence. This region contains the other. Adjacent to the terminal palindrome is an AT-rich region that is determined. There are two orientations of the 242-base-pair terminal of the tandem repeat sequence. Similarly, the related parvovirus H-3 the implications of these results are discussed.

Record Date Created: 19820910

6/7/51 (Item 51 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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Incomplete genomes of the parvovirus minute virus of mice: selective conservation of genome termini, including the origin for DNA replication. Faust E A; Ward D C

Journal of virology (UNITED STATES) Oct 1979, 32 (1) p276-92, ISSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

Record type: Completed

mice viral DNA. Type II D-DNAs were virus-specific, double-stranded hairpin designated as type I D-DNA and type II D-DNA, could be distinguished on the roughly equal amounts. Type I D-DNAs were predominantly single-stranded, ecombinant molecules in which the self-complementary sequences derived population, sequences which map between coordinates 47.3 and 87.1 were from the total D-DNA population during serial undiluted passage, suggesting high-multiplicity passage was heterogeneous, ranging in size from 15 to 70% from both genomic termini were conserved. The 5' terminus was modified sequence and consisted of sequences derived almost exclusively from within analyzed. The DNA isolated from incomplete virions derived from a single oopulation lacked between 90 and 95% of the internal wild-type genome sequences which mapped entirely at the 5' end of the viral genome between 5.0 map units (250 nucleotides) at both ends of the viral genome. Moreover, molecules whose complementary strands were covalently continuous at that these molecules are not competent for DNA Replication but arise as the these miniature recombinant molecules were selectively amplified during positions 85.0 and 100. Furthermore, type II molecules were gradually lost clearly underrepresented. However, the extent and position of the deletions future studies on parvovirus DNA replication, transcription, and cell-virus isolated, and the incomplete viral genomes contained therein have been the wild-type genome sequence was seen in the total type I D-DNA in individual molecules varied significantly. The shortest molecules in the high-multiplicity passage and after serial undiluted passage have been serial undiluted passage and were therefore believed to contain all of the approximately, 2,000 nucleotides. Two distinct types of molecules, basis of their degree of secondary structure, and these were present in contrast to the type I genomes, these hairpin molecules consisted of here for minute virus of mice should be valuable generally as aids to result of fatal replication errors. Deletion mutants of the type described relative to the analogous wild-type structure. Although virtually all of critical recognition sites necessary for the replication of minute virus of Deletion mutants of minute virus of mice arising during a single of the intact viral genome, with an average molecular length of variable sites distal to the 5' end of the viral minus strand. In sharp

Record Date Created: 19800616

6/7/52 (Item 52 from file: 155)

DIALOG(R)File 155:MEDLINE(R)

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Nucleotide sequence of the self-priming 3' terminus of the single-stranded DNA extracted from the parvovirus Kilham rat virus. Salzman L A; Fabisch P Journal of virology (UNITED STATES) Jun 1979, 30 (3) p946-50, ISSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Languages: ENGLISH Main Citation Owner: NLM Record type: Completed The parvovirus genome is a linear, single-stranded DNA molecule with double-stranded hairpin termini. The 3' terminus can serve in vitro as a self-primer for the synthesis of a double-stranded viral DNA intermediate. We have sequenced the nucleotides in the 3' terminus and propose a model for the secondary structure of the terminus and the in vitro origin of	 E2 1 ARVIN E3 0 *ARVO? OR AAV OR ADENOASSOCIAT? OR ADENO(W)ASSO E4 1 ARVR E5 1 ARV2 E6 2 ARW E7 1 ARW-7 E8 11 ARX E9 1 ARXIDA
replication for the complementary viral DNA strand. Record Date Created: 19791128	3(7)
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\$15.82 Estimated cost this search \$29.74 Estimated total session cost 4.248 DialUnits	S2 4872 TERMINUS OR TERMINI OR TELOMER? S3 45 S1 AND S2
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nucleic acid (I) and a CARE-DRI. BIOTECHNOLOGY - Preferred Nucleic Acid and cap genes operably linked to a CARE sequence with a CARE-DRI; (8) a adenoviral DNA-Binding Protein (Ad DBP), the gene of the Ad DBP, and following contacting the cell with a CARE-DRI. (I) Comprises a CARE and cell, comprising contacting the cell with a CARE-DRI (the DNA sequence Sequences: In (I) the CARE comprises a defined nucleotide sequence (N1) Herpes viruses, the adenoviral DNA-Binding Protein (Ad DBP), the gene recombinant AAV preparations, comprising contacting cells harboring rep comprising a CARE, and contacting the cell-line with a CARE-DRI; and Comprises retroviral Long Terminal Repeats (LTRs). Preferred Methods: papilloma virus selected from HeLa, Hela32, SIHA, CASKI cells and cells a polynucleotide sequence heterologous to AAV, a poly-linker comprising operably linked to a DNA sequence heterologous to AAV and to the cells method (IX) of producing recombinant AAV preparation, comprising and (b) an integrated copy of an MV-derived vector, comprising a DNA derived from them. In (III) the CARE-DRI is selected from Adenoviruses, for the amplification of a DNA sequence operably linked to a CARE and sequence operably linked to a CARE and integrated into the genome of a sequence of interest flanked by AAV Inverted Terminal Repeats (ITRs) of the Ad DBP, and any gene transfer vector expressing the Ad DBP. In (IV) the DNA sequence to be amplified further encodes the rep genes of cell-lines (V) and/or (VI). In the methods (VIII) and (IX) the CARE-DRI with a CARE-DRI; (3) a method (IV) for the amplification of a DNA (4) a highly producing rAAV packaging cell-line (V) comprising: (a) an CARE-DRI); (5) a highly producing rAAV packaging cell-line (VI) ransfecting the cell-line (VI) with a plasmid harboring a rAAV genome inked to a CARE sequence; and (b) a second integrated copy of the cap to be amplified encodes the cap genes of an Adeno-Associated Virus); mutant/fragment still promotes the amplification of a DNA sequence gene; (6) a cell-line (VII) comprising an integrated CARE sequence integrated copy of the rep and cap genes, operably linked to a CARE; comprising: (a) an integrated copy of the rep and cap genes, operably (9) a kit (X) for amplifying a DNA sequence in a cell, comprising the from which the cell-line is derived; (7) a method (VIII) of producing integrated into the genome of a cell and operably linked to the CARE an Adeno Associated Virus. In the method (VII) the cells are the integrated into the genome of a cell, comprising contacting the cell (replication of the integrated rep and cap genes is inducible by a is selected from Adenoviruses, Herpes viruses (preferred), the given in the specification (or a fragment/mutant, provided the several cloning sites and/or genetic elements from a virus. (I) In (II) and (III) the cell is a cell-line harboring part of human AUTHOR: SALVETTI A; CHADEUF G; TESSIER J; MOULLIER P; LINDEN M R; ABSTRACT: DERWENT ABSTRACT: NOVELTY - An isolated nucleic acid sequence PATENT NUMBER: WO 200246359 PATENT DATE: 20020613 WPI ACCESSION element (CARE) from an Adeno-Associated Virus (AAV), and a second DNA NATIONAL APPLIC. NO.: WO 2001EP15418 APPLIC. DATE: 20011206 contacted with a CARE-dependent replication inducer (CARE-DRI) comprising a first DNA sequence comprising a cis-acting replication sequence operably linked to the CARE, is new Amplification of the Cis-acting replication elements from an Adeno-Associated Virus (AAV), isolated nucleic acid sequence occurs when the isolated nucleic acid PRIORITY APPLIC. NO.: US 251576 APPLIC. DATE: 20001207 sequence is integrated in the genome of a cell and the cell is adeno-associated virus production by packaging cell culture with useful for producing cell lines that express AAVs - recombinant 1542 PARVO? OR AAV OR ADENOASSOCIAT? OR 4872 TERMINUS OR TERMINI OR TELOMER? 0302115 DBR Accession No.: 2003-03900 PATENT (c) 2003 Thomson Derwent & ISI. All rts. reserv. PATENT ASSIGNEE: UNIV NANTES 2002 1772 ORIGIN (2N)REPLICATION DIALOG(R)File 357:Derwent Biotech Res. potential application in gene therapy 2114 LEFT OR RIGHT 2002-706808 (200276) 55 S5 NOT S3 ADENO(W)ASSOCIAT? 46 S1 AND S7 Items Description 45 S1 AND S2 58 S1 AND S4 55 S5 NOT S3 LANGUAGE: English ?ts8/7/14181927 P; EPSTEIN A L

1772 S7 1542 SI

?s s1 and s7

S8

CARE-DRI is a herpesvirus mutant selected from DELTAICPO, HP66, HR94,

any gene transfer vector expressing the Ad DBP. In particular, the

DETAILED DESCRIPTION - INDEPENDENT CLAIMS are also included for: (1) a

comprising: (a) operably linking a DNA sequence to an isolated CARE;

method (II) for the amplification of a DNA sequence in a cell,

(b) introducing the sequence operably linked to the CARE into the cell

genome; and (c) contacting the cell with a CARE-DRI; (2) a method (III)

comprises a CARE sequence, in sense or antisense orientation. The CARE

inked to the integrated rep and cap genes is in sense orientation, and

and 1178ts. Preferred Cell-Lines: In (V) the AAV-derived vector

of packaging cell lines that express recombinant AAV. ADVANTAGE - The from Adenoviruses, Herpes viruses, the adenoviral DNA-Binding Protein is operably linked to a CARE sequence. (V) And (VI) are derived from a comprising Adenoviruses, Herpes viruses and Retroviruses. The kit may integrated copy of the cap gene operably linked to a CARE sequence. In Repeats (LTRs). Preferred Kits: In the kit (X) the CARE-DRI is selected methodologies. USE - The nucleic acid (I) may be used in the production human cell-line harboring part of human papilloma virus such as HeLa, HeLa32, SIHA and CASKI cells. In the cell-lines (V) - (VII) 1 or more orientation. The packaging cell-line (V) may further comprises a second sequence (I) may be produced via standard recombinant and synthetic the packaging cell-line (VI) the second integrated copy of the cap gene expressing the Ad DBP. The kit further comprises a rep expression the CARE comprised in the integrated rAAV vector is in antisense further comprises a purified Rep protein. Preparation: The nucleic acid nucleic acid (I), derived from the genome of AAV-2 behaves like a (Ad DBP), the gene of the Ad DBP, and any gene transfer vector replication origin in the presence of AAV Rep proteins and a helper cassette enclosed in a plasmid or in a vector selected from the group of the integrated elements is flanked by retroviral Long Terminal virus (adenovirus). EXAMPLE - No example given (76 pages)

8/1/8

DIALOG(R)File 357:Derwent Biotech Res.

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0297472 DBR Accession No.: 2002-19319

A cis-acting element that directs circular adeno-associated virus replication and packaging - adeno-associated virus replication and

packaging, expression in cell culture use in gene therapy AUTHOR: MUSATOV S, ROBERTS J, PFAFF D, KAPLITT M

CORPORATE AFFILIATE: Cornell Univ Rockefeller Univ

CORPORATE SOURCE. Kaplitt M, Cornell Univ, Weill Med Coll, Dept Neurosurg, 525 E 68 St, New York, NY 10021 USA

IOURNAL: JOURNAL OF VIROLOGY (76, 24, 12792-12802) 2002

ISSN: 0022-538X

LANGUAGE: English

ABSTRACT: AUTHOR ABSTRACT - A novel pathway of adeno-associated virus

replication marked by the assembly of circular monomer duplex intermediates (cAAV) has been recently discovered. In the present report we identify a single AD domain of the inverted terminal repeat as a minimal origin of cAAV replication. A small internal palindrome (BB'), necessary for optimal Rep-inverted terminal repeat interaction, does not contribute to the efficiency of cAAV replication, whilethe terminal resolution site is an essential cis-acting element. Furthermore, recombinant cAAV vectors that encompass only the AD domain replicate exclusively in a circular form and no detectable linear

duplex replicative intermediates are generated, suggesting that both pathways of AAV replicationare independent and can be separated. In addition, we show that cAAVs areefficient templates for encapsidation of single-stranded DNA genomes, an observation that assigns a biological role for these novel replication species. Together, these findings shed new light on the current model of AAV replication and packaging. (11 pages)

81/1/8

DIALOG(R)File 357:Derwent Biotech Res.

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0238719 DBR Accession No.: 99-08820 PATENT

Transcriptionally-activated adeno-associated virus inverted terminal repeat

- recombinant adeno-associated virus vector-mediated chloramphenicol-acetyltransferase gene transfer and expression in human

bronchial epithelium cell culture for gene therapy AUTHOR: Feldhaus A L

CORPORATE SOURCE: Seattle, WA, USA.

PATENT ASSIGNEE: Targeted-Genet. 1999

PATENT NUMBER: WO 9920773 PATENT DATE: 990429 WPI ACCESSION NO.: 99-288312 (9924)

PRIORITY APPLIC. NO.: US 955400 APPLIC. DATE: 971021

NATIONAL APPLIC. NO.: WO 98US21937 APPLIC. DATE: 981020

LANGUAGE: English

ABSTRACT: A polynucleotide (PN) containing a transcriptionally-activated useful for the production of improved recombinant AAV vectors which may PN; a mammalian cell containing any PN; and a method of packaging a recombinant AAV vector. The transcriptionally-activated ITRs may be of cystic fibrosis gene therapy. In an example, recombinant AAV vectors adeno-associated virus (AAV) inverted terminal repeat (ITR) (less than especially the cystic fibrosis transmembrane conductance regulator gene containing the chloramphenicol-acetyltransferase (EC-2.3.1.28) gene, transcriptionally-activated ITR; an AAV virus particle containing any wild-type ITR, a transcriptionally-activated ITR, a D sequence, a trs or a portion of a wild-type ITR, a plasmid containing a PN and an element selected from the group of a replication origin and a reporter transcriptionally-activated ITR and a second ITR chosen from a 400 bp in length) is new. Also claimed are: a PN consisting a were used to transfect human bronchial epithelial IB3 cells. (55pp) gene; a PN, further consisting a gene operably linked to a be particularly useful for the packaging of large transgenes,

8/1/19

DIALOG(R)File 357: Derwent Biotech Res.

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0236812 DBR Accession No.: 99-06913 PATENT

New autonomous parvo virus-based vector - recombinant virus vector-mediated

S10 148756 3 OR LEFT S11 674 S1 AND S10 S10 148756 3 OR LEFT ADENO(W)ASSOCIAT? 58 S1 AND S4 55 S5 NOT S3 848 DUPLICAT? Items Description 45 S1 AND S2 12 S12(3N)S11 46 S1 AND S7 ? s s12 (3n)s11 and s1 15420 TWO 1595 LEFT 148756 S10 ? s two or duplicat? 16166 S12 674 S11 148115 3 1542 S1 ? s s1 and s10 ?s 3 or left PATENT NUMBER: WO 9911802 PATENT DATE: 990311 WPI ACCESSION NO.: PATENT NUMBER: WO 9745550 PATENT DATE: 971204 WPI ACCESSION NO.: ABSTRACT: A new DNA sequence has an adeno virus-5 (Ad5) inverted terminal ABSTRACT: An autonomous parvo virus-based vector (I) (minute virus of mice) AUTHOR: Zhang W W; Alemany R; Dai Y; Josephs S; Balague C; Ayares D; PATENT ASSIGNEE: Yeda-Res.Develop.; Univ.Negev-Ben-Gurion 1999 desired specific target DNA, is new. The target DNA is a chromosomal or heterologous DNA transfer and expression in Escherichia coli for use in Mini-adenoviral vectors carrying the minimal cis-element of the adenoviral genome - adeno virus vector construction in packaging cell culture, and binding and nicking sites recognized by an autonomous parvo virus recombination. The vector contained a binding site recognized by the integration of the heterologous DNA into the host cell. The vector may episomal site of a host cell, consisting of a DNA sequence containing NATIONAL APPLIC. NO.: WO 97US10218 APPLIC. DATE: 970530 for (I) directed integration of a heterologous DNA and a method for which is capable of stably integrating a heterologous DNA into a regulatory protein, the replication origin (ori) of (I), a DNA sequence encoding the regulatory protein and a heterologous DNA to be regulatory protein for the initiation of localized DNA synthesis and repeat (ITR), a packaging signal, a transcription control region, an which contains a sequence based on/related to the 3' or 5' or and integrated. Also claimed is a 53 bp oligonucleotide (II) (specified) be useful for the correction of a genetic defect, for the prevention or NATIONAL APPLIC. NO.: WO 98IL419 APPLIC. DATE: 980828 inserted into a host cell (e.g. Escherichia coli), becoming a target identifying sites for site-specific chromosomal and episomal human cancer and infectious disease prevention and gene therapy PRIORITY APPLIC, NO.: US 791218 APPLIC, DATE: 970131 green fluorescent protein or human Factor-VIII expression in a PRIORITY APPLIC. NO.: IL 121676 APPLIC. DATE: 970901 reatment of cancer, or infectious diseases in humans. (45pp) CORPORATE SOURCE: Rehovot, Italy; Beer Sheva, Israel. 0221156 DBR Accession No.: 98-02753 PATENT transgenic mouse model for gene therapy testing (c) 2003 Thomson Derwent & ISI. All rts. reserv. CORPORATE SOURCE: Deerfield, IL, USA. AUTHOR: Winocour E; Tal J; Corsini J A DIALOG(R)File 357: Derwent Biotech Res. PATENT ASSIGNEE: Baxter 1997 LANGUAGE: English LANGUAGE: English 99-205195 (9917) 98-032656 (9803) Schneiderman R

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pCMV-hFVIII-mini, AFP-pEGFP-1, mAFP-hFVIII/pGKNeo and RIP-pEGFP-from-BS
                                                     alpha-fetoprotein Alb-E5 AFP-3 or EBB14 gene homologous recombination
                                                                                                                                                                                                                                                                                                    of the DNA does not encode an Ad protein. A human telomere, a human
                                                                                                                                                                              numan or SV40 virus replication origin, alphoid DNA, T-antigen or oriP
                                                                                                                                                                                                                                           and Epstein-Barr virus nuclear antigen-1) sequence, where the remainder
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          protein, and the effector gene may encode human Factor-VIII. The vector
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         transgenic mouse by microinjection into embryonic stem cells. Plasmid
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              vectors are new. The transgenic mouse may be used to test gene therapy
                                                                                                                 arm, or an adeno-associated virus ITR) or episomal maintenance (e.g. a
                                                                                                                                                                                                                                                                                                                                                                                                                             may also be present. The reporter gene may encode green fluorescent
                                                                                                                                                                                                                                                                                                                                                                 albumin or alpha-1-antitrypsin promoter and a liver-specific enhancer
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            replication-defective Ad vector. The vector may be used to generate a
effector or reporter gene, and a genomic integration (e.g. a albumin or
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               may be packaged in a cell culture to give an infectious
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            1542 PARVO? OR AAV OR ADENOASSOCIAT? OR
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     4872 TERMINUS OR TERMINI OR TELOMER?
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              S12 16166 TWO OR DUPLICAT?
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               strategies using the vectors. (193pp)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     >>>Warning: unmatched quote found
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       2114 LEFT OR RIGHT
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1542 SI

S13 12 S12 (3N)S11 AND S1

? t s13/kwic/1 13/KWIC/1 DIALOG(R)File 357:(c) 2003 Thomson Derwent & ISI. All rts. reserv.

- ...vector, by introducing vector having nucleotide sequence, adenovirus inverted terminal repeats and packaging sequence, and adeno associated virus terminal repeat, into cell, and culturing cell virus vector expression in host cell use...
- ..ABSTRACT: terminal repeats of adenovirus flanking NS, adenovirus packaging sequence linked to inverted terminal repeat, and adenoassociated virus terminal repeat linked to 3' end of NS, into cell expressing adenovirus early gene...
 - ..flanking NS, adenovirus packaging sequence linked to one of the inverted terminal repeats, and an adeno associated virus terminal repeat sequence operably linked to the 3' end of NS, where the first vector lacks a second adeno associated virus terminal repeat sequence, and lacks one or more adenovirus early gene region such as...
 - .. gene which is lacking from the first vector; and (iii) optionally a helper adenovirus or adeno associated virus; (b) introducing the first vector, and optionally the genome of a helper adenovirus or adeno associated virus, into the cell to produce a transformed cell; and (c) culturing the transformed cell...
- ...to produce a second vector chosen from: (i) a third vector comprising in operable combination adeno associated virus terminal repeat DD sequence, first and second inverted copies of NS flanking the adeno associated virus terminal repeat-DD sequence, left and right inverted terminal repeats of adenovirus flanking the...
 - ... lacks adenovirus E3 early gene region, or by providing the first recombinant vector that comprises adeno associated virus rep gene region, introducing first vector into the cell, and culturing the transformed cell...
- ... rep proteins and to produce a second vector. WIDER DISCLOSURE (1) recombinant vectors including adenovirus/ adeno associated virus vectors and mini-adenovirus (mAd) vectors; (2) cells containing the vectors of (1); and...
- ...as E1, E2, and E4 gene region, when the method involves the first vector comprising adeno associated virus rep gene region. ACTIVITY Immunostimulant; Antianemic; Antilipemic; Nootropic; Cytostatic; Dermatological. No biological data is...
- and dimeric mini-adenoviruses (mAd) in exemplary 293 cells were performed as follows. Recombinant adenovirus/ adeno associated virus (Ad/ AAV) hybrid virus had the left end of Ad5 containing the inverted terminal repeat (ITR) and packaging domain, the AAV TR D sequence, an green fluorescent protein (EGFP)/neomycin (Neo) expression cassette from the plasmid pTRUF2, an intact AAV terminal repeat with a double D sequence (TR-DD), and the remainder of the Ad...

- mini-adenoviruses using the parental Ad/ AAV EGFP/Neo virus, 293 cells which complement the E1 deletion in the hybrid virus to allow virus replication were infected with a cellular lysate containing the parental Ad/ AAV hybrid virus. Two days after infection, cleared cellular lysates were prepared and treated with DNase I and RNase A. Ad/ AAV and mAd viruses were separated on a CsCl2 step gradient. The lower band represented full...
- ... resistant, confirming that it was packaged within the virions. During normal replication of wild type AAV with an Ad helper virus, both monomer length as well as dimer length AAV genome products were observed as part of the replication pathway. DNA analysis was also carried out on the viral particles, which demonstrated that full virus particles contained the parental Ad/ AAV hybrid virus genome as a single DNA molecule 36 kbase in size. The E/M...
- ... DNA (nucleotides 1-420) containing the Ads ITR and packaging domain, as well as the AAV TR D sequence. The remainder of the AAV terminal repeat was missing from the mAd genome. The dimeric form contained a duplicated monomer genome where the left end of Ad5 (nucleotides 1-420), AAV TR D sequence and the EGFP-Neo expression cassette were duplicated in an inverted manner. An intact AAV TR was present at the junction of the duplication. (191 pages)
 - DESCRIPTORS: recombinant adeno virus, adeno associated virus vector-mediated gene transfer expression in mouse, human host cell, inverted terminal repeat, packaging...
- familial hypercholesterolemia, Lesch-Nyhan syndrome, phenylalanine hydroxylas phenylketonuria, muscular dystrophy, cystic fibrosis therapy, gene therapy parvo virus mammal animal immunostimulant antianemic antipemic nootropic cytostatic (22, 07)

? s s12(3n)s10 and s1

16166 S12

148756 S10

317 S12(3N)S10

1542 SI

S14 3 S12(3N)S10 AND S1

? t s14/kwic/1-3

14/KWIC/1

DIALOG(R)File 357:(c) 2003 Thomson Derwent & ISI. All rts. reserv. ...ABSTRACT: at undetectable levels at day 1 after injection. By day 35, expression had increased by two logs and was 3 - 4 fold higher than the levels of CAT from pCFI-CAT at this time point...

... hybrid promoter is also useful in the context of viral vectors, e.g. adenovirus and adeno - associated vectors, and in other tissues such as muscle or brain. ADVANTAGE - (V) confers high and...

14/KWIC/2

DIALOG(R)File 357:(c) 2003 Thomson Derwent & ISI. All rts. reserv.

encoding the hFIX gene in rhesus macaques - vector-mediated gene liver-targeted delivery of recombinant adeno - associated virus Sustained high-level expression of human factor IX (hFIX) after transfer and expression in...

.ABSTRACT: using hyperimmune serum from a rhesus monkey that had received an adenoviral vector encoding hFIX. Two macaques having 3 and 40 rAAV genome equivalents/cell, respectively, in liver tissue had 4% and

DESCRIPTORS: adeno - associated virus vector-mediated human Factor-IX, cytomegalo virus enhancer, beta-actin promoter gene transfer,

mammal animal blood-clotting protein herpes virus embryo kidney DNA polymerasechain reaction, SDS-PAGE, ELISA, antibody, Western blot hybridization analysis, appl. hemophilia B gene therapy parvo virus amplification analysis immunoassay DNA... expression.

Mutational analysis of the adeno - associated virus type 2 (AAV2) capsid DIALOG(R)File 357:(c) 2003 Thomson Derwent & ISI. All rts. reserv. gene and construction of AAV2 vectors with altered tropism

ABSTRACT: To obtain a comprehensive genetic map of the adeno - associated capsid gene were constructed by site-directed mutagenesis. Several virus capsid gene, 93 mutants at 59 different positions in the AAV types of mutants were studied including...

particle (VP)-3. Two of these mutants were insertions at the N and ... in what were likely to be beta-barrel in the capsid protein virus C termini of VP3, suggesting that ...

.. ligand in the N-terminal regions of VP1 or VP2 could change the tropism of AAV. The results provided information on AAV capsid functional domains and are useful for future design of AAV vectors for targeting of specific tissues. (51 ref)

DESCRIPTORS: adeno - associated virus vector construction, site-directed mutagenesis, capsid gene, altered tropism, appl. gene therapy parvo virus (Vol. 19, No. 22)

Set Items Description

1542 PARVO? OR AAV OR ADENOASSOCIAT? OR

ADENO(W)ASSOCIAT?

4872 TERMINUS OR TERMINI OR TELOMER?

2114 LEFT OR RIGHT 45 SI AND S2

58 S1 AND S4

55 S5 NOT S3

1772 ORIGIN (2N)REPLICATION 46 SI AND S7

148756 3 OR LEFT

05mar03 11:56:29 User208669 Session D2223.5 \$1.00 4 Type(s) in Format 95 (KWIC) \$0.00 116 Type(s) in Format 6 \$38.79 2.153 DialUnits File357 \$16.15 5 Type(s) in Format 7 16166 TWO OR DUPLICAT? 12 S12 (3N)S11 AND S1 3 S12(3N)S10 AND S1 674 S1 AND S10 ? log hold **S14 S13 S12**

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\$55.94 Estimated cost File357

\$3.02 TELNET

\$58.96 Estimated cost this search

\$88.70 Estimated total session cost 6.401 DialUnits

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Connecting via Winsock to Dialog

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Reconnected in file 357 05mar03 12:14:12

* * New CURRENT Year ranges installed

File 357:Derwent Biotech Res. _1982-2003/Mar W2

*File 357: File is now current. See HELP NEWS 357 (c) 2003 Thomson Derwent & ISI

Alert feature enhanced for multiple files, etc. See HELP ALERT.

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Cost is in DialUnits

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\$1.50 0.083 DialUnits File357

\$1.50 Estimated cost File357

\$0.01 TELNET

\$1.51 Estimated cost this search

\$1.51 Estimated total session cost 0.083 DialUnits

File 155:MEDLINE(R) 1966-2003/Mar W1 (c) format only 2003 The Dialog Corp.

Set Items Description

? s aav or adenoassociat? or adeno(w)associat? 1075 AAV

S4 7894 NICK **S**2 element. Furthermore, recombinant cAAV vectors that encompass only the AD S1 1704 AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT? assembly of circular monomer duplex intermediates (cAAV) has been recently A novel pathway of adeno-associated virus (AAV) replication marked by the single-stranded DNA genomes, an observation that assigns a biological role Weill Medical College of Cornell University, 525 East 68th Street, New addition, we show that cAAVs are efficient templates for encapsidation of Laboratory of Neurobiology and Behavior, The Rockefeller University, inverted terminal repeat as a minimal origin of cAAV replication. A small pathways of AAV replication are independent and can be separated. In internal palindrome (BB'), necessary for optimal Rep-inverted terminal discovered. In the present report we identify a single AD domain of the domain replicate exclusively in a circular form and no detectable linear for these novel replication species. Together, these findings shed new Journal of virology (United States) Dec 2002, 76 (24) p12792-802, duplex replicative intermediates are generated, suggesting that both repeat interaction, does not contribute to the efficiency of cAAV A cis-acting element that directs circular adeno-associated virus replication, while the terminal resolution site is an essential cis-acting Musatov Sergei, Roberts Jill, Pfaff Donald, Kaplitt Michael, et al ight on the current model of AAV replication and packaging. (c) format only 2003 The Dialog Corp. All rts. reserv. SSN 0022-538X Journal Code: 0113724 4198590 22326831 PMID: 12438604 1540 ADENO(W)ASSOCIAT? 80 MINIMAL (W)ORIGIN DIALOG(R)File 155:MEDLINE(R) 43 ADENOASSOCIAT? Record Date Created: 20021119 Document type: Journal Article 1115086 ASSOCIAT? Main Citation Owner: NLM eplication and packaging. 99279 MINIMAL Record type: Completed 2 SI AND S2 137442 ORIGIN Languages: ENGLISH York, NY 10021, USA. 2502 ADENO ? s minimal (w)origin 1704 S1 80 S2 ?s sl and s2 ? t s3/7/1 **S**5

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mediate vectorial unwinding of the DNA duplex via an ATP-dependent helicase
                                          1704 AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT?
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      The Rep78 gene product of adeno-associated virus (AAV) self-associates to
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                amino acid sequences required for maximal self-association occur between
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Molecular Hematology Branch, National Heart, Lung, and Blood Institute,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               experiments with a bacterially expressed maltose-binding protein-Rep78
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       The Rep78 and Rep68 proteins of adeno-associated virus (AAV) are
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       studies with N- and C-terminal truncation mutant forms of Rep indicate that
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      hrough the use of a mammalian two-hybrid system. Rep-Rep protein
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               replication. Genetic and biochemical studies have identified Rep mutants
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    hat demonstrate a trans-dominant negative phenotype in vitro and in vivo,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         synthesized in a coupled in vitro transcription-translation system. Mapping
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 fusion protein in combination with [358]methionine-labeled Rep78
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          esidues 164 and 484. Site-directed mutagenesis identified two essential
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           activity, thus initiating a strand displacement mechanism of viral DNA
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    vitro and in vivo. Self-association of Rep78 in vivo was demonstrated
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       interaction was confirmed in vitro through communoprecipitation
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Journal of virology (UNITED STATES) Jun 1997, 71 (6) p4461-71,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               notifs within this 321-amino-acid region: (i) a putative alpha-helix
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         suggesting the possibility that multimerization of Rep is essential for
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               replication initiator proteins that bind the viral replicative-form origin
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 of replication, nick the origin in a site- and strand-specific fashion, and
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           certain replicative functions. In this study, we have investigated the
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ability of the largest of the Rep proteins, Rep78, to self-associate in
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           form a hexameric complex in the presence of AAV ori sequences
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      (c) format only 2003 The Dialog Corp. All rts. reserv.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ISSN 0022-538X Journal Code: 0113724
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                                                                                                   80 MINIMAL (W)ORIGIN
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                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Bethesda, Maryland 20892, USA.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Document type: Journal Article
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tems Description
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cross-linking experiments indicated that Rep78 forms a hexameric complex in bearing a 3,4-hydrophobic heptad repeat reminiscent of those found in inphosphate-binding motif. Deletion of either of these regions from the full-length polypeptide resulted in severe impairment of Rep-Rep coiled-coil domains and (ii) a previously recognized nucleoside interaction. In addition, gel filtration chromatography and protein the presence of AAV ori sequences.

Record Date Created: 19970609

DIALOG(R)File 155:MEDLINE(R)

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Sequence requirements for stable binding and function of Rep68 on the adeno-associated virus type 2 inverted terminal repeats.

Molecular Hematology Branch, National Heart, Lung, and Blood Institute, Chiorini J A; Wiener S M; Owens R A; Kyostio S R; Kotin R M; Safer B Bethesda, MD 20892-1654. Journal of virology (UNITED STATES) Nov 1994, 68 (11) p7448-57,

SSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Main Citation Owner: NLM Languages: ENGLISH

Record type: Completed

Replication of the palindromic inverted terminal repeats (ITRs) of adeno-associated virus type 2 requires several functions of the viral

protein (MBP)-Rep68 delta produced in Escherichia coli and wild-type (wt) report demonstrates the ability of both recombinant fusion maltose-binding nonstructural Rep proteins. These include binding to the ITR, nicking of site (trs), and then strand displacement and synthesis from the nick. This Rep68 to bind to a linear truncated form of the ITR, delta 57 ITR, with the double-stranded replication intermediate at the terminal resolution similar affinity as to the wthairpin ITR. A dissociation constant for

MBP-Rep68 delta of approximately 8 x 10(-10) M was determined for the wt ITR and delta 57 ITR probes. Truncation of delta 57 ITR to generate delta 28 ITR, which retains the GCTC repeat motif but not the trs, bound at least nonspecific sequence restored the ability of MBP-Rep68 delta to bind to 10 times less efficiently than delta 57 ITR. Extension of delta 28 ITR with delta 28 ITR. Thus, high-affinity binding would appear to require

spanned the region protected by MBP-Rep78 in DNase I footprinting. The binding activity of either MBP-Rep68 delta or wt Rep68 to hairpin ITR or Cleavage of the delta 57 ITR probe with Ddel, which truncates the flanking inhibited the binding of MBP-Rep68 delta. The requirements for stable binding were further defined with a series of oligonucleotide probes which sequence and was previously shown to inhibit binding by Rep68, also stabilization by flanking sequence as well as the intact GCTC repeat motif. delta 57 ITR was indistinguishable. However, the binding activity of

MBP-Rep68 delta to DNA does not appear to correlate with trs endonuclease nonhairpin delta 57 ITR was approximately 100-fold less efficient than its activity. The nicking and covalent linkage of MBP-Rep68 delta to the stabilization of MBP-Rep68 delta binding, its presence does affect the trs inkage to a hairpin-containing ITR. Therefore, although the hairpin portion of the ITR does not appear to play a role in recognition and cleavage activity of the protein.

Record Date Created: 19941117

DIALOG(R)File 155:MEDLINE(R)

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Adeno-associated virus DNA replication in vitro: activation by a maltose binding protein/Rep 68 fusion protein.

Ward P; Urcelay E; Kotin R; Safer B; Berns K I

Department of Microbiology, Hearst Microbiology Research Center, Cornell

Journal of virology (UNITED STATES) Sep 1994, 68 (9) p6029-37, University Medical College, New York, New York 10021

SSN 0022-538X Journal Code: 0113724

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well as the AAV DNA; however, linear pBR322 DNA was not replicated. When plasmid construct containing intact AAV DNA. When the recombinant plasmid observed when the template was either linear double-stranded AAV DNA or a construct was used as the template, there was replication of pBR322 DNA as The adeno-associated virus (AAV) nonstructural protein Rep 68 is required sequences or the newly synthesized sequence hairpinned, switched template strands, and replicated the AAV DNA. Replication was linear for 4 h, during addition of Rep 68 to an extract from uninfected HeLa cells supports AAV which time 70% of the maximal synthesis took place. An additional finding the intact plasmid and led to separation of the AAV sequences from those of the vector, a process which has been termed rescue. There was no evidence a site-specific nick 124 nucleotides from the 3' end of AAV DNA; the site proceeding toward the 3' terminus. When the template was the plasmid for viral DNA replication. An in vitro assay has been developed in which replication process when a fusion of the maltose binding protein and Rep 68, expressed in Escherichia coli, was used in the assay. Replication was that replication could initiate on the products of rescue. Rep 68 can make the plasmid construct was the template, replication appeared to initiate on of the nick has been called the terminal resolution site. Our data are most construct, either elongation continued past the junction into pBR322 DNA replication. In this paper, we report characterization of the consistent with initiation occurring at the terminal resolution site and

was that the Rep fusion could resolve AAV dimer length duplex intermediates into monomer duplexes without DNA synthesis.

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Features of the adeno-associated virus origin involved in substrate ecognition by the viral Rep protein.

Snyder R O; Im D S; Ni T; Xiao X; Samulski R J; Muzyczka N

Department of Microbiology, State University of New York at Stony Brook Medical School 11794-8621. Journal of virology (UNITED STATES) Oct 1993, 67 (10) p6096-104,

SSN 0022-538X Journal Code: 0113724

Contract/Grant No.: 5 PO1 CA2814607; CA; NCI; AI25530; AI; NIAID; RO1

GM3572302; GM; NIGMS

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We previously demonstrated that the adeno-associated virus (AAV) Rep68

and Rep78 proteins are able to nick the AAV origin of DNA replication at the terminal resolution site (trs) in an ATP-dependent manner. Using four types of modified or mutant substrates, we now have investigated the

substrate requirements of Rep68 in the trs endonuclease reaction. In the first kind of substrate, portions of the hairpinned AAV terminal repeat

internal palindromes of the AAV terminal repeat were active in the were deleted. Only deletions that retained virtually all of the small

biochemical evidence that the secondary structure of the terminal repeat endonuclease reaction. This result confirmed previous genetic and

genome. The mutant was nicked at a 50-fold-lower frequency relative to a substrate, the trs was moved eight bases further away from the end of the was an important feature for substrate recognition. In the second type of

wild-type origin, and the nick occurred at the correct trs sequence despite secondary structure. It also suggested that the minimum trs recognition sequence extended three bases from the cut site in the 3' direction. The ts new position. This finding indicated that the endonuclease reaction required a specific sequence at the trs in addition to the correct

mismatch substrates contained a wild-type sequence on the strand normally mismatch mutants were capable of being nicked in the presence of ATP cut but an incorrect sequence on the complementary strand. All of the third type of substrate harbored mismatched base pairs at the trs. The However, there was substantial variation in the level of activity,

suggesting that the sequence on the opposite strand may also be recognized during nicking. Analysis of the mismatch mutants also suggested that a single-stranded trs was a viable substrate for the enzyme. This

he enzyme. We concluded that substrate recognition by the AAV Rep protein involves at least two and possibly as many as four features of the AAV cleaved efficiently by the enzyme provided that the correct strand was wild-type did. This finding indicated that the sequence at the cut site was not involved in recognition of the terminal repeat for specific binding by nterpretation was confirmed by analysis of the fourth type of substrate tested, which contained a single-stranded trs. This substrate was also substrates with mutant trss bound the Rep protein as efficiently as the present in the substrate. In addition, the single-stranded substrate no onger required ATP as a cofactor for nicking. Finally, all of the terminal repeat. (ABSTRACT TRUNCATED AT 400 WORDS)

Record Date Created: 19931012

Items Description

1704 AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT?

80 MINIMAL (W)ORIGIN

2 SI AND S2

7894 NICK

12 S1 AND S4

? log hold

05mar03 12:22:14 User208669 Session D2223.7

\$0.00 14 Type(s) in Format 6 \$2.85 0.891 DialUnits File155

\$1.05 5 Type(s) in Format 7

\$1.05 19 Types

\$3.90 Estimated cost File155

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Items Description Set

1704 AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT?

80 MINIMAL (W)ORIGIN **S**2

2 SI AND S2

full-length pA2Y1 DNA molecules was increased by MNNG treatment of cells in DNA primase, and DNA topoisomerase I. Furthermore, restriction endonuclease N-methyl-N'-nitro-N-nitrosoguanidine (MNNG). A pUC19-based plasmid, a dose-dependent manner. In addition, DNA synthesis of plasmid pA2Y1 was studied in vitro. Extracts derived from MNNG-treated CHO-9 and L1210 cells designated pA2Y1, which contains the left terminal repeat sequences (TRs) representing the AAV origin of replication and the p5 and p19 promoter but suggested that the reaction is largely dependent on DNA polymerase alpha, origin of AAV type 2 DNA replication cloned into a plasmid is sufficient to cells. Following transfection of plasmid pA2Y1 or plasmid pUC19 as a control, density labeling by a bromodeoxyuridine and DpnI resistance assay displayed greater synthesis of DpnI-resistant full-length pA2Y1 molecules (adeno-associated virus) can be induced by a variety of genotoxic agents in than did nontreated controls. Experiments with specific enzyme inhibitors Institut fur Virusforschung/Angewandte Tumorvirologie, Deutsches the absence of coinfecting helper virus. Here we investigated whether the napping analysis of the in vitro reaction products revealed the occurrence activity and to allow selective DNA amplification in carcinogen-treated origin-containing plasmid. Furthermore, the amount of DpnI-resistant Journal of virology (UNITED STATES) Jun 1991, 65 (6) p3175-84, DNA amplification of the helper-dependent parvovirus AAV suggested a semi-conservative mode of replication of the AAV acks any functional parvoviral genes is shown to confer replication Origin of adeno-associated virus DNA replication is a target of Krebsforschungszentrum, Heidelberg, Federal Republic of Germany. promote replication activity in cells treated by the carcinogen (c) format only 2003 The Dialog Corp. All rts. reserv. S6 1829 ORIGIN (1W) REPLICATION Yalkinoglu A O; Zentgraf H; Hubscher U SSN 0022-538X Journal Code: 0113724 16924727 91237836 PMID: 2033669 carcinogen-inducible DNA amplification. DIALOG(R)File 155:MEDLINE(R) Document type: Journal Article 96349 REPLICATION Main Citation Owner: NLM ? s origin (1w) replication S7 12 S1 AND S6 Record type: Completed 12 SI AND S4 137442 ORIGIN Languages: ENGLISH 1704 SI 1829 S6 ?ssl and s6 ? t s7/7/12

of specific initiation at the AAV origin of DNA replication. Though elongation was not very extensive, extracts from carcinogen-treated cells markedly amplified the AAV origin region. Our results, including electron microscopic examination, suggest that the AAV origin/terminal repeat structure is recognized by the cellular DNA replicative machinery induced or modulated by carcinogen treatment in the absence of parvoviral gene products.

Record Date Created: 19910626

?ts7/7/1111

1/1/1

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Adeno-associated virus (AAV) site-specific recombination does not require a Rep-dependent origin of replication within the AAV terminal repeat.

Young S M; Samulski R J

Curriculum in Genetics and Molecular Biology, Gene Therapy Center,

University of North Carolina, Chapel Hill, NC 27599, USA.

Proceedings of the National Academy of Sciences of the United States of America (United States) Nov 20 2001, 98 (24) p13525-30, ISSN 0027-8424 Journal Code: 7505876

Contract/Grant No.: DK51880; DK; NIDDK; HL 48347; HL; NHLBI

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required for AAV DNA replication are located on chromosome (ch) 19. Both extensive rearrangement and amplification of ch-19 sequences independent of of targeted integration in human cells. An AAV Rep binding element (RBE) Adeno-associated virus (AAV) is the only known eukaryotic virus capable terminal repeat (ITR) cis-acting sequences in targeted integration an AAV cotransfection in the presence or absence of Rep. Our results demonstrated genomes were unable to rescue and replicate. In addition, Rep78 induced prerequisite for site-specific recombination and suggests AAV targeting is similar frequency. Molecular characterization of the mutant ITR integrants mediated by Rep78/68-dependent replication from the ch-19 origin of that, in the presence of Rep78, both ITR substrates targeted to ch-19 with ested. Complementation analysis confirmed that the mutant targeted viral rs mutant incapable of supporting viral replication was tested. Wild-type ch-19 RBE and trs elements have been shown to be essential for viral wild-type or mutant targeting substrate. These studies demonstrate that confirmed the presence of the trs mutation in the majority of samples targeting to this locus. To characterize the role of the AAV inverted Rep-dependent nicking of the viral cis-acting trs sequence is not a eplication (ori). These studies have significant impact toward the and mutant substrates were assayed for targeted integration after and terminal resolution site (trs) identical to the viral terminal repeats

understanding of AAV site-specific recombination and the development of targeting vectors.

Record Date Created: 20011121

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Analysis of the terminal repeat binding abilities of mutant

ideno-associated virus replication proteins.

Yang Q; Trempe J P

Department of Biochemistry and Molecular Biology, Medical College of

Ohio, Toledo 43699-0008.

Journal of virology (UNITED STATES) Jul 1993, 67 (7) p4442-7, ISSN

3022-538X Journal Code: 0113724

Document type: Journal Article

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Main Citation Owner: NLM

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The adeno-associated virus (AAV) Rep78 and Rep68 proteins play essential roles in viral DNA replication, trans activation of viral gene expression,

and suppression of oncogene-mediated cellular transformation. By using an gene, we mapped the regions of the Rep78 protein that mediate binding to extensive set of linker insertion and deletion mutations in the replication

codons 25 to 62, 88 to 113, 125 to 256, and 346 to 400 abolished binding. he AAV origin of replication in vitro. Deletions that removed amino acid

Alterations in several other regions of the protein affected the binding affinity of the mutant proteins. All of the mutant proteins that support

AAV DNA replication or p40 trans activation bound to the terminal repeat Several mutant rep genes that failed to suppress oncogene-mediated cellular sequence, thus verifying the importance of binding for these functions.

ransformation produced proteins that were capable of binding to the AAV

Record Date Created: 19930712 terminal repeat sequences

? s trs or terminal(w)resolution(w)(site or sites)

769 TRS

191235 TERMINAL

88924 RESOLUTION

336559 SITE

364178 SITES

781 TRS OR TERMINAL (W) RESOLUTION (W) (SITE OR SITES) 33 TERMINAL(W)RESOLUTION(W)(SITE OR SITES) 8 8

Items Description

1704 AAV OR ADENOASSOCIAT? OR ADENO(W)ASSOCIAT? SI

80 MINIMAL (W)ORIGIN 2 SI AND S2 S2 S3

12 S1 AND S4 7894 NICK S4 S5 S6 S7

1829 ORIGIN (1W) REPLICATION

12 S1 AND S6

781 TRS OR TERMINAL (W) RESOLUTION (W) (SITE OR SITES)

ssl and s8

1704 S1

781 S8

39 S1 AND S8 S₀

? t s9/7/23

9/7/23

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Minimum origin requirements for linear duplex AAV DNA replication in

Ward P; Berns K I

Department of Microbiology, W. R. Hearst Microbiology Research Center,

Cornell University Medical College, New York, New York 10021, USA.

Virology (UNITED STATES) Jun 1 1995, 209 (2) p692-5, ISSN 0042-6822 lournal Code: 0110674

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Document type: Journal Article Languages: ENGLISH

Main Citation Owner: NLM

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virus (AAV) Rep 68/78 protein. When a linear duplex template was used, initiation of one round of DNA replication was achieved when nucleotides We have investigated the minimal requirements for a functional origin of DNA replication in an in vitro assay which requires the adeno-associated required. Deletion of an additional 11 nucleotides from the terminal repeat -56 of the AAV inverted terminal repeat (in the flop orientation) were present at the terminus of the template. A terminal resolution site was not olocked the reaction.

Record Date Created: 19950711

DIALOG(R)File 155:MEDLINE(R)

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Adeno-associated virus (AAV) type 5 Rep protein cleaves a unique terminal resolution site compared with other AAV serotypes.

Chiorini J A; Afione S; Kotin R M

Molecular Hematology Branch, National Heart, Lung, and Blood Institute, Bethesda, Maryland 20892, USA.

Journal of virology (UNITED STATES) May 1999, 73 (5) p4293-8, ISSN 0022-538X Journal Code: 0113724

Document type: Journal Article

Main Citation Owner: NLM Languages: ENGLISH

Record type: Completed

sequences. AAV2 Rep78 cleaved only a type 2 ITR DNA sequence, and AAV5 proteins cannot cross-complement in the packaging of a genome with an AAV2 ITRs of other AAV serotypes. Comparison of the TRSs in the AAV2 ITR, the AAV5 ITR, and the AAV chromosome 19 integration locus identified some Rep78 cleaved only a type 5 probe efficiently. Mapping of the AAV5 ITR TRS Adeno-associated virus (AAV) replication depends on two viral components distinct virus compared to the other cloned AAV serotypes. Whereas the Rep nverted terminal repeat (ITR) sequences in cis. AAV type 5 (AAV5) is a identified a distinct cleavage site (AGTG TGGC) which is absent from the proteins and ITRs of other serotypes are interchangeable and can be used to conserved nucleotides downstream of the cleavage site but little homology evel of replication instead of at viral assembly. AAV2 and AAV5 Rep AAV5 ITR; however, comparison of terminal resolution site (TRS) endonuclease activities showed a difference in specificity for the two DNA produce recombinant viral particles of a different serotype, AAV5 Rep binding activities demonstrate similar affinities for either an AAV2 or or replication: the AAV nonstructural proteins (Rep) in trans, and TR. In vitro replication assays indicated that the block occurs at the

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A novel terminal resolution-like site in the adeno-associated virus type

2 genome.

Wang X S; Srivastava A

Department of Medicine, Walther Oncology Center, Indiana University

School of Medicine, Indianapolis 46202, USA.

Journal of virology (UNITED STATES) Feb 1997, 71 (2) p1140-6, ISSN 0022-538X Journal Code: 0113724

Contract/Grant No.: DK-49218; DK; NIDDK; HL-48342; HL; NHLBI; HL-53586;

H. NHLBI

Document type: Journal Article

Languages: ENGLISH

Main Citation Owner: NLM

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The adeno-associated virus 2 (AAV) contains a single-stranded DNA genome of which the terminal 145 nucleotides are palindromic and form T-shaped important role in AAV DNA replication and resolution, since each of the hairpin structures. These inverted terminal repeats (ITRs) play an TRs contains a terminal resolution site (trs) that is the target site for

AAV plasmids and the Rep-mediated resolution of the ITRs during AAV DNA Rep-mediated excision of the viral genome during rescue from recombinant replication, we constructed recombinant AAV genomes that lacked either the infected cells or from recombinant AAV plasmids. To distinguish between with the AAV DNA sequences that lie outside the ITRs, and the ITRs also rescue and replication of the AAV genome occurred from these plasmids he AAV rep gene products (Rep). However, the Rep proteins also interact the vector sequences. These studies suggest that the Rep-mediated cleavage the AAV promoters at map unit 5 (AAV p5) that also contains an RBS. the left end of the AAV genome. This site was precisely mapped to one of eft or the right ITR sequence and one of the Rep-binding sites (RBSs). No sequences was clearly detected from the plasmid that lacked the AAV left following transfection into adenovirus type 2-infected human KB cells, as Furthermore, deletion of this RBS abolished the rescue and replication of expected. However, excision and abundant replication of the vector at the RBS during viral DNA replication may, in part, account for the TR, suggesting the existence of an additional putative excision site in play a crucial role in excision of the proviral genome from latently generation of the AAV defective interfering particles.

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Sequence requirements for binding of Rep68 to the adeno-associated virus erminal repeats.

Ryan J H; Zolotukhin S; Muzyczka N

Department of Molecular Genetics and Microbiology, College of Medicine, Jniversity of Florida, Gainesville 32610, USA.

Journal of virology (UNITED STATES) Mar 1996, 70 (3) p1542-53,

Contract/Grant No.: HL/DK 50257; HL; NHLBI; PO1 CA2814607; CA; NCI; RO1 SSN 0022-538X Journal Code: 0113724

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GM3572302; GM; NIGMS

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22-bp sequence (CAGTGAGCGAGCGAGCGCAG) in which substitutions which contained the Rep binding element (RBE) within the A stem of the TR. protein. Mutagenesis experiments were done with a 43-bp oligonucleotide Experiments in which two adjacent base pairs of the RBE were substituted We have used reciprocal competition binding experiments with mutant define the sequences within the adeno-associated virus (AAV) terminal simultaneously with nucleotides that produced transversions identified a repeat (TR) that are involved in site-specific binding to the AAV Rep substrates and chemical modification interference assays to precisely

neasurably

the greatest effect. Dinucleotide mutations within 18 (GTGAGCGAGCGAGC) of threefold. Dinucleotide mutations within a 10-bp core sequence (GCGAGCGAGC) esidues within the sequence CTTTG. This sequence was present in the same hydrazine were performed on both the linear A-stem sequence and the entire affected the binding affinity. Although the 22-bp RBE contains the GAGC suggested that the remaining bases within the CTTTG motif as well as other bases within the B and C palindromes make contacts with the Rep protein, analysis suggested that the A-stem RBE contains only a single Rep binding secondary structure element which consists of the B and C palindromes, and assays on the linear A stem identified the 18-bp sequence described above suggest that the GAGC motifs alone are not the only sequences specifically are three distinct elements within the hairpinned AAV TR that contribute to recognized by Rep. The effects of substitutions within the 22-bp sequence to binding, chemical modification experiments using dimethyl sulfate and AAV TR in both the flip and flop hairpinned configurations. Interference were found to decrease binding affinity by more than 10-fold. Single-base were relatively symmetrical, with nucleotides at the periphery of the RBE hairpinned TR substrates also identified the 18-bp sequence as important substitutions within the 10-bp core sequence lowered the binding affinity by variable amounts (up to fivefold). The results of the mutagenesis the mutant analysis and to determine the relative contribution of each base A-stem binding element. Interference assays also allowed us to search for contribute to binding. The largest effect was seen by modification of two T with hairpinned substrates and found that linear substrates bound Rep less for binding. However, the interference patterns on the two strands within residues within the small internal palindromes of the TR (B and C) that motifs that have been found in all known Rep binding sites, our results as essential for binding. G, C, and T residues on both strands contributed the RBE and the relative contributions of the individual bases to binding efficiently. Our results were consistent with our previous model that there site rather than two or more independent sites. To confirm the results of internal palindrome furthest away from the trs. We also determined the having the least effect on binding affinity and those in the middle having of the mutagenesis experiments. Interference assays with complete elative binding affinity of linear substrates containing a complete RBE were clearly different between the hairpinned substrates and the linear albeit with lower affinities. Regardless of whether the TR was in the flip or flop orientation, most of the contact points were clustered in the small to binding, and the interference pattern correlated well with the results oinding affinity or to efficient nicking at the trs: the A-stem RBE, the the 22 bp were found to decrease the binding affinity by at least and flop orientations of the TR. In addition, the interference pattern position relative to the terminal resolution site (trs) in both the flip

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Sequence requirements for stable binding and function of Rep68 on the adeno-associated virus type 2 inverted terminal repeats.

Molecular Hematology Branch, National Heart, Lung, and Blood Institute, Chiorini J A; Wiener S M; Owens R A; Kyostio S R; Kotin R M; Safer B Bethesda, MD 20892-1654.

Journal of virology (UNITED STATES) Nov 1994, 68 (11) p7448-57, SSN 0022-538X Journal Code: 0113724

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protein (MBP)-Rep68 delta produced in Escherichia coli and wild-type (wt) report demonstrates the ability of both recombinant fusion maltose-binding nonstructural Rep proteins. These include binding to the ITR, nicking of site (trs), and then strand displacement and synthesis from the nick. This Rep68 to bind to a linear truncated form of the ITR, delta 57 ITR, with the double-stranded replication intermediate at the terminal resolution Replication of the palindromic inverted terminal repeats (ITRs) of adeno-associated virus type 2 requires several functions of the viral similar affinity as to the wthairpin ITR. A dissociation constant for

MBP-Rep68 delta of approximately 8 x 10(-10) M was determined for the wt MBP-Rep68 delta to DNA does not appear to correlate with trs endonuclease spanned the region protected by MBP-Rep78 in DNase I footprinting. The binding activity of either MBP-Rep68 delta or wt Rep68 to hairpin ITR or Cleavage of the delta 57 ITR probe with Ddel, which truncates the flanking ITR and delta 57 ITR probes. Truncation of delta 57 ITR to generate delta 28 ITR, which retains the GCTC repeat motif but not the trs, bound at least nhibited the binding of MBP-Rep68 delta. The requirements for stable nonspecific sequence restored the ability of MBP-Rep68 delta to bind to sequence and was previously shown to inhibit binding by Rep68, also binding were further defined with a series of oligonucleotide probes which 10 times less efficiently than delta 57 ITR. Extension of delta 28 ITR with stabilization by flanking sequence as well as the intact GCTC repeat motif. nonhairpin delta 57 ITR was approximately 100-fold less efficient than its stabilization of MBP-Rep68 delta binding, its presence does affect the trs activity. The nicking and covalent linkage of MBP-Rep68 delta to the delta 57 ITR was indistinguishable. However, the binding activity of inkage to a hairpin-containing ITR. Therefore, although the hairpin delta 28 ITR. Thus, high-affinity binding would appear to require portion of the ITR does not appear to play a role in recognition and cleavage activity of the protein.

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Interaction of the adeno-associated virus Rep protein with a sequence within the A palindrome of the viral terminal repeat.

McCarty D M; Ryan J H; Zolotukhin S; Zhou X; Muzyczka N

Department of Microbiology, School of Medicine, University at Stony

Brook, New York 11794.

Journal of virology (UNITED STATES) Aug 1994, 68 (8) p4998-5006,

SSN 0022-538X Journal Code: 0113724

Contract/Grant No.: PO1 CA2814607, CA; NCI; RO1 GM3572302, GM; NIGMS, T32

AI25530; AI; NIAID

Document type: Journal Article

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We have characterized a Rep binding sequence which is within the A stem region of the adeno-associated virus terminal repeat (TR) and compared its affinity with that of the complete hairpinned TR for pure Rep68. Both the A stem and the complete TR substrates produced a complex pattern of protein-DNA complexes in which at least six different bound species could be distinguished. Competition experiments suggested that the dissociation constant for the A stem sequence is approximately 125-fold higher than that for the complete TR. The competition experiments also suggested that the average number of Rep molecules per TR substrate molecule under conditions

of saturating substrate is 3.7:1, while for the A stem substrate, the ratio

the A stem complex rather than the actual number of Rep molecules per DNA substrate from the protein-DNA complexes was approximately fourfold faster the 25-bp A stem region. Both of these mutants completely abolished binding the protein-DNA complexes with the two kinds of substrates, suggesting that Muzyczka, J. Virol. 68:4988-4997, 1994) as essential for binding. Each of these mutants eliminated some but not all of the repeating GAGC motifs in sequence of the A stem that had been identified in the accompanying report other mutants produced seven- or five-base substitutions within the 25-bp substrates, however, were complex, suggesting that substrate was being substitution mutants within the A stem of the TR. A five-base mutant near complexes, no major difference was seen in the mobility or the pattern of released from at least two different kinds of protein-DNA complexes at (D. M. McCarty, D. J. Pereira, I. Zolotukhin, X. Zhou, J. H. Ryan, and N. is 10:1. In spite of the apparent difference in protein-to-DNA ratio in the the difference in protein-to-DNA ratio was due to the lower stability of molecule. At least some of the difference in stability of the two kinds of complexes was due to the fact that the dissociation rate of the A stem than that of the complete TR. The dissociation rate curves for both the terminal resolution site (trs site) had little effect on binding. Two different rates. In addition, we have analyzed binding to several

to the A stem substrate but only partially reduced binding in the context of the complete hairpinned TR. Furthermore, neither mutant altered the pattern of Rep-DNA complexes produced.(ABSTRACT TRUNCATED AT 400 WORDS)

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Identification of linear DNA sequences that specifically bind the

adeno-associated virus Rep protein.

McCarty D M; Pereira D J; Zolotukhin I; Zhou X; Ryan J H; Muzyczka N

Department of Microbiology, School of Medicine, University at Stony

Brook, New York 11794.

Journal of virology (UNITED STATES) Aug 1994, 68 (8) p4988-97,

SSN 0022-538X Journal Code: 0113724

Contract/Grant No.: PO1 CA2814607; CA; NCI; RO1 GM3572302; GM; NIGMS; T32

AI25530; AI; NIAID

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within a 25-bp sequence of the A stem of the adeno-associated virus (AAV) homogeneity to reexamine the binding properties of the Rep protein. We find Rep78. The ability to recognize the linear DNA sequence within the A stem terminal repeat proximal to the B and C palindromes. This has been shown suggests a mechanism by which the Rep protein could act as a repressor or a DNA affinity column containing the 25-bp sequence can be used to purify that Rep68 is capable of binding to a linear DNA sequence that is contained computer analysis suggests that sequences similar to the A stem element are provides a mechanism by which the Rep protein can be oriented on the present within the three AAV promoter regions. Electrophoretic mobility shift experiments clearly demonstrate that the p5 promoter contains a Rep initiator sequence and the TATA binding site. This position immediately We have used baculovirus-expressed Rep68 that has been purified to binding the A stem recognition element, as demonstrated by the fact that a binding sequence. DNase protection experiments indicate that the Rep synthetic oligonucleotide containing the 25-bp region in the absence of the binding sequence within the p5 promoter is located between the YY1 other sequences within the terminal repeat. Rep78 was also capable of transactivator of p5 transcription by interacting with either YY1 or TBP. conclusively by demonstrating that Rep68 could specifically bind to a In addition, gel shift experiments suggest that the p19 promoter also terminal repeat so that only the correct strand is cut at the terminal resolution site (trs site) during terminal resolution. In addition,

contains a Rep binding site. The presence of Rep binding sites upstream of

any apparent secondary structure. (ABSTRACT TRUNCATED AT 400 WORDS) heterologous Rep binding sequence within pBR322 DNA. A comparison of the the accompanying report (D. M. McCarty, J. H. Ryan, S. Zolutukhin, X. Zhou, and N. Muzyczka, J. Virol. 68:4998-5006, 1994), we examine the relative sequences within the A stem, p5, and pBR322 binding sites suggests that a Finally, we also have reexamined the ability of Rep68 and Rep78 to cut at repeating GAGC motif is at least part of the Rep recognition sequence. In both promoters suggests that these sites may be involved in coordinate the trs site in substrates that do not contain the B and C palindromes or regulation of AAV transcription. In addition, we have identified a affinity of Rep to the A stem site and the complete terminal repeat. Record Date Created: 19940815

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In vitro replication of adeno-associated virus DNA.

Ni T H; Zhou X; McCarty D M; Zolotukhin I; Muzyczka N

Department of Microbiology, State University of New York, Stony Brook Medical School 11794.

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termini, a key step in AAV DNA replication. We have now cloned the AAV developed an efficient in vitro AAV DNA replication system which mimics the in vivo behavior of AAV in every respect. With no-end AAV DNA as the AAV nonstructural proteins, Rep78 and -68, are site-specific endonucleases Adeno-associated virus (AAV) is distinct from previously reported systems and DNA helicases that are capable of resolving covalently closed AAV permissive replication. In previous work, we demonstrated that two of the expression system. Using the baculovirus-expressed proteins, we have synthesis and requires coinfection with adenovirus to establish completely nonstructural proteins Rep78, Rep68, and Rep52 in the baculovirus The study of eukaryotic viral DNA replication in vitro has led to the starting substrate, the reaction required an adenovirus-infected cell identification of cellular enzymes involved in DNA replication. in that it is believed to replicate entirely by leading-strand DNA

not support DNA replication. A mutant in the AAV terminal resolution site

(trs) was defective for DNA replication in the in vitro assay. Little, if

any, product was formed in the absence of the adenovirus-infected HeLa cell

extract and the presence of either Rep78 or Rep68. Rep52, as expected, did

heavy-heavy product DNA in the presence of bromo-dUTP when examined on CsCl virus 40 in vitro system. Replication of the complete AAV DNA molecule was supporting AAV DNA replication than adenovirus-infected extracts. Thus, the demonstrated by the following criteria (i) Most of the monomer and dimer vitro. The reduced ability of uninfected HeLa extracts to support complete 32P-deoxynucleoside triphosphate could be detected in DpnI-resistant monomer replicative form and was linear for at least 4 h after the lag. The DNA replication was not due to a defect in terminal resolution but rather rate of incorporation in the reaction was comparable to that in the simian product DNAs were completely resistant to digestion with DpnI. (ii) requirement for adenovirus infection in vivo was partially duplicated in to a defect in the reinitiation reaction or in elongation. Rep78 produced a characteristic monomer-dimer pattern of replicative intermediates, but surprisingly, Rep68 produced little, if any, dimer replicative form. The Virtually all of the starting substrate was converted to heavy-light or reaction had a significant lag (30 min) before incorporation of extract. In general, uninfected HeLa extracts were less efficient in density gradients. (ABSTRACT TRUNCATED AT 400 WORDS)

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Features of the adeno-associated virus origin involved in substrate

Snyder R O, Im D S, Ni T; Xiao X; Samulski R J; Muzyczka N ecognition by the viral Rep protein.

Department of Microbiology, State University of New York at Stony Brook Medical School 11794-8621.

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We previously demonstrated that the adeno-associated virus (AAV) Rep68 and Rep78 proteins are able to nick the AAV origin of DNA replication at the terminal resolution site (trs) in an ATP-dependent manner. Using four types of modified or mutant substrates, we now have investigated the substrate requirements of Rep68 in the trs endonuclease reaction. In the biochemical evidence that the secondary structure of the terminal repeat internal palindromes of the AAV terminal repeat were active in the first kind of substrate, portions of the hairpinned AAV terminal repeat endonuclease reaction. This result confirmed previous genetic and were deleted. Only deletions that retained virtually all of the small

he enzyme. We concluded that substrate recognition by the AAV Rep protein mismatch substrates contained a wild-type sequence on the strand normally genome. The mutant was nicked at a 50-fold-lower frequency relative to a suggesting that the sequence on the opposite strand may also be recognized nismatch mutants were capable of being nicked in the presence of ATP substrate, the trs was moved eight bases further away from the end of the during nicking. Analysis of the mismatch mutants also suggested that a wild-type origin, and the nick occurred at the correct trs sequence despite cut but an incorrect sequence on the complementary strand. All of the was an important feature for substrate recognition. In the second type of involves at least two and possibly as many as four features of the AAV secondary structure. It also suggested that the minimum trs recognition sequence extended three bases from the cut site in the 3' direction. The third type of substrate harbored mismatched base pairs at the trs. The cleaved efficiently by the enzyme provided that the correct strand was not involved in recognition of the terminal repeat for specific binding by wild-type did. This finding indicated that the sequence at the cut site was interpretation was confirmed by analysis of the fourth type of substrate ts new position. This finding indicated that the endonuclease reaction ested, which contained a single-stranded trs. This substrate was also substrates with mutant trss bound the Rep protein as efficiently as the present in the substrate. In addition, the single-stranded substrate no single-stranded trs was a viable substrate for the enzyme. This However, there was substantial variation in the level of activity, onger required ATP as a cofactor for nicking. Finally, all of the required a specific sequence at the trs in addition to the correct terminal repeat.(ABSTRACT TRUNCATED AT 400 WORDS)

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